APPENDIX 1



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## (54) LITHIUM ION SECONDARY BATTERY

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## ABSTRACT

A lithium ion secondary battery includes a positive electrode capable of absorbing and desorbing lithium ion, a negative electrode capable of absorbing and desorbing lithium ion, a porous film interposed between the positive electrode and the negative electrode, and a non-aqueous electrolyte, the porous film being adhered to a surface of at least the negative electrode. The porous film includes an inorganic filler and a first binder: The content of the first binder in the porous film is 1.5 to 8 parts by weight per 100 parts by weight of the filler. The first binder includes a lirst rubber including an acrylonitrile unit: The first rubber is waterinsoluble and has a decomposition temperature of 250° C. or higher. The negative electrode includes a negative electrode active material capable of absorbing and desorbing lithium ion and a second binder, and the second binder includes a second rubber particle and a water-soluble polymer.

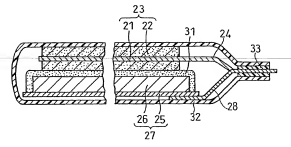
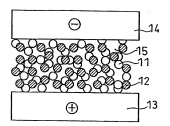


FIG. 1



F | G. 2

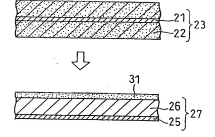


FIG. 3

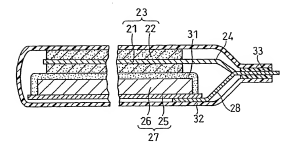


FIG. 4

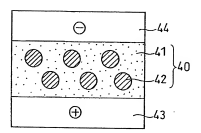
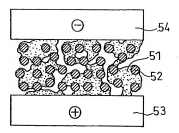


FIG. 5



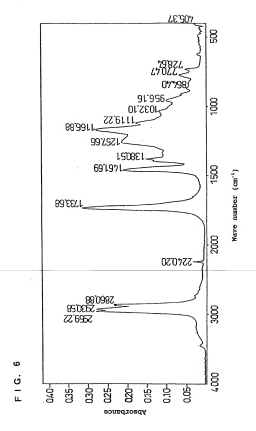


FIG. ,7

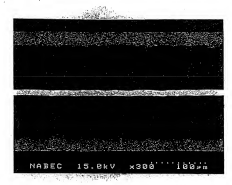
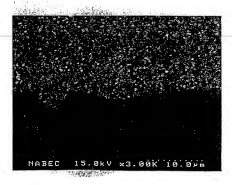


FIG. 8



## LITHIUM ION SECONDARY BATTERY

#### TECHNICAL FIELD

[0001] The present invention relates to a lithium ion secondary battery having a porous film adhered to a surface of at least a negative electrode, the porous film comprising a filter and a hinder. The thitium ion secondary hattery of the present invention has a low occurrence rate of a short circuit and is excellent in safety in terms of heat resistance and the like

#### BACKGROUND ART

[9002] As electronic devices are becoming more and more portable and wireless, small and lightweight lithium ion secondary hatteries with a higher energy density are gaining attention as the power source for these devices. Lithium ion secondary hatteries have a positive electrode comprising a lithium-containing transition metal oxide or the like, a negative electrode comprising a carbon material or the like, a separator interposed between the positive electrode and the negative electrode, and a non-aqueous electrolyte.

[0003] For the separator of lithium ion secondary batteries, a drawn resin film (a sheet separator) is used generally. For the raw material of the resin film, polyolefins such as polyothylene and polypothylene are used, However, as in polyothylene and polypothylene and polypothylene and polypothylene and in a seposed to a heisthrikage is caused when the film is exposed to a heisthrikage is caused when the film is exposed to a heisthrikage is caused when the film is exposed to the battery steep of resin film may cause deterioration of over 150° C. the shrinkage of resin film may cause deterioration diplex such to a nail penetrates the hattery (at the time of nail penetration test), a short circuit portion expands due to a short circuit reaction heat which is generated instantly, causing further reaction beat, rising a possibility of an acceleration of shormal overheat.

[9094] As schematically shown in FIG. 4, there has been proposed a usage of a paste electrolyse 40 for the function of a separator. The paste electrolyte 40 includes a great amount of liquid electrolyte 41 including a thickener, and an electrically insulating filler 42. The filler 42 functions as a spacer between a positive electrode 43 department of the pastern of the pastern between a positive electroled 43 and a negative electrode 44 (Japanese Laid-Open Pattern Publication No. 1hei 10-5578 liquid electrolyte in which a viscosity is enhanced by the thickener and of an electrically insulating filler. Injuid electrolyte is with a viscosity is enhanced by the thickener and of an electrically insulating filler. Injuid electrolyte is sufficiently included therein. Therefore, the paste electrolyte is excellent in that a certain level of fillumin on conductivity can be secured. I not ever, there are defects that the strength electron of the pastern of the p

[0005] Further, there has been proposed a technology to form a portus film including a filler comprising inorganic particles on a surface of a sheet separator comprising a resin film (Japanese Laid-Open Patent Publication No. 2001-319634, and Japanese Laid-Open Patent Publication No. 2001-319634, and Japanese Laid-Open Patent Publication No. 2002-37301, therever, since the provis film is fromed on the surface of the sheet separator in these proposals, there are decies that the provisa film artinas as the sheet separator shrinks. These techniques are aiming at suppression of a strike of the shrinks of the shrinks of the shrinks of the shrinks. These techniques are aiming at suppression of a shrinks. These techniques are aiming at suppression of a shrinks that the shrinks of the shrinks of the shrinks that the shrinks of the shrinks o [9006] On the other hand, there has been proposed a technique in which a prous fill no comprising a resin with a low glass transition temperature is formed on an electrod-(Dapanese Laid-Open Patent Publication No. Hei 11-144406). This proposal is intended to develop the shitdown effect by softening the resin with a low glass transition temperature when a heat is generated due to a short circuit, in such proposal, for example in an all pentertain test, due to the generated heat at the time of an internal short circuit, the temperature locally exceeds several handred "C. depending on conditions, causing excessive softening or bouring of resint. As a result, provise slift adjoints, and may conclude the conditions of the condition of the condition of the conditions of the condition o

[0007] There has been also proposed a technique to form a protective buyer comprising inorganic particles such as alumina or the like, and a water-soluble polymer on an electrode (Japanese Lidó-Open Paten Publication No. 15c) 9-147916. For the water-soluble polymer, a polyacyfic acid derivative, a cellulous edurativite, and the like are used. Based on this proposal, since the protective layer includes inorganic particles with excellent bear resistance, suppression of the deformation of the protective layer itself at the time of heat generation due to short circuit can be experient on the control of the protective layer itself at the time of heat generation due to short circuit can be experient on the control of the protective layer is the first the control of the protective layer is the protective layer to the

[0008] However, currently, for a negative electrode of lithium ion secondary batteries, styrene-butadiene copolymer (SBR) or rubber particles comprising a modified product thereof are generally used as a negative electrode binder in amay cases. His is because in the case of further particles, only a small usage amount is necessary, compared with the case of polythyridene fluoride (PDP) conventionally use as a negative electrode binder, and lithium ion acceptability of negative electrode improves.

[9000] In the case where rubber particles are to be included in a negative electrode material mixture, such as this charter comprising water-soluble polymer should be used to together with the rubber particles. For the water should polymer, cellulose resin is the mainstream material. When posteric help reincluding, the water-soluble polymer is applied on such negative electrode, the thickners in the reagtive electrode swell by water included in the protective layer before drying, enssing a deformation of the negative electrode schedulor when the control water layer before drying, enssing a deformation of the negative electrode. Although the negative electrode iccuracy-ended the electrode schedulor has provided to a practical use, the yield drops dramatically.

[0010] Also, there has been examined a separator provided by forming a thin film of paste including a resin binder dissolved in a solvent, and a filler on a surface of a positive electrode or a negative electrode, and then drying the formed film (Japanese Laid-Open Patent Publication No. Hei 10-106530). In such paste, fluorocarbon resin, polyolefin resin, or the like is included as the resin binder. Such separator is excellent in that a certain level of strength is secured. However, when the resin binder deposits on the surfaces of filler particles after dissolved in a solvent, as schematically shown in FIG. 5, the areas of filler particles 52 covered with a resin binder 51 become large. As a result, gaps among filler particles decrease while strength increases, causing an insufficient moving path between a positive electrode 53 and a negative electrode 54 for the liquid electrolyte or lithium ion to move. Additionally,

although many techniques have been proposed to form a paste thin film including a resin binder dissolved in a solvent and a filler on a surface of a positive electrode or a negative electrode, the same problems will occur (Japanese Laid-Open Patent Publication No. Hei 7-220759, Patent Publication No. 3371301, Patent Publication No. 34262531.

## DISCLOSURE OF INVENTION

[0011] The present invention aims to improve the safety of a lithium ion secondary battery, while preventing a deformation of a negative electrode having a higher lithium ion acceptability, by adhering a porous film excellent in heat resistance to a surface of at least the negative electrode.

[0012] That is, the present invention relates to a lithium ion secondary battery comprising a positive electrode capable of absorbing and desorbing lithium ion, a negative electrode capable of absorbing and desorbing lithium ion, a porous film interposed between the positive electrode and negative electrode, and a non-aqueous electrolyte: wherein the porous film is adhered to a surface of at least the negative electrode; the porous film comprises an inorganic filler and a first binder; a content of the first binder in the porous film is 1.5 to 8 parts by weight per 100 parts by weight of the inorganic filler; the first binder comprises a first rubber including an acrylonitrile unit; the first rubber is waterinsoluble and has a decomposition temperature of 250° C. or higher; the negative electrode comprises a negative electrode active material capable of absorbing and desorbing lithium ion and a second binder; and the second binder includes a second rubber particle and a water-soluble poly-

[9013] A stacked-type battery can be obtained when the positive electrode and the negative electrode are stacked with the porous film interposed therebetween, and a woundtype battery can be obtained when the positive electrode and the negative electrode are sprainly wound with the porous film interposed therebetween. The present invention is applicable to any type of the battery.

[9014] Since the prous film is quite excellent in bear resistance and is being althred to an electrode suffice, the porous film does not shrink by heat, rendering the electrontion by softening or burning difficult. Therefore, a safe lithium non secondary battery can be obtained. Also, since the first binder included in the prorus (film is water-insoluble, drops in yields due to the deformation of the negative electrode can be avoided, even when the prorus film is combined with a high performance negative electrode including a thicknew comprising a water-soluble polymer. It is desirable that the thickness of the porous film is 0.5 µm or more and 20 µm or less.

[0015] In the porous film, it is desirable that the first binder has no crystalline melting point or has a crystalline melting point of 250° C. or higher. It is desirable that the first nubber comprises core-shell type particles and has an adhesive surface portion. Also, it is desirable that the first nubber includes a polyacytomitrike chain.

[0016] It is desirable that in the negative electrode, the water-soluble polymer includes a methylcellulose unit. It is desirable that the second rubber particle includes a styrene unit and a butadrene unit.

[0017] The present invention also aims to effectively prevent a short circuit of lithium ion secondary battery by optimizing the balance between lithium ion acceptability of negative electrode and lithium ion permeability of porous film.

[0018] That is, it is desirable that a content of the second binder in the negative electrode is 1.5 to 3 parts by weight of the negative electrode active material. Within this range, the balance between lithium ion acceptability of negative electrode and fifthium ion permeability of porous film can be maintained at optimal condition.

[0019] When the negative electrode includes a great amount of the binder, the surface of the negative electrode active material is covered with the binder for the most part, to decrease lithium acceptability of the negative electrode. When lithium ion acceptability of negative electrode declines in a battery having no porous film, metallic lithium deposits at gaps between the negative electrode and the sheet separator. The drawbacks involved with such state are developed as an increase in irreversible capacity over a long period of time. On the other hand, when the porous film is attached on a surface of the negative electrode, since there is no gap between the negative electrode and the separator, metallic lithium deposits in the porous film. As a result, a defect occurs as a short circuit over a short period of time. By setting the content of the second binder in the negative electrode to 1.5 to 3 parts by weight per 100 parts by weight of the negative electrode active material, such defect can be suppressed.

[0020] Additionally, by optimizing the balance of lithium acceptability of negative electrode and lithium ion permeability of porous film, a lithium ion secondary battery excellent in high-rate characteristic can be obtained, in addition to suppressing the occurrence of defects.

[0021] The present invention also aims to achieve a higher safety by using a combination of a porous film excellent in heat resistance and a conventional sheet separator.

[0022] That is, by further providing a separator interposed between the positive electrode and the negative electrode to a lithium ion secondary battery of the present invention, a higher safety can be achieved. It is preferable that the thickness of the separator is 8 mm or more and 30 mm or less.

[0023] The present invention also aims to achieve a higher yield by controlling the surface roughness of the porous film.

[0024] That is, it is preferable that the surface roughness of the porous film is smaller than the surface roughness of an electrode surface to which the porous film is addired, in view of preventing a separation of a material mixture from the electrode and a decline in yield. By making the surface roughness on the electrode surface small, a separation of the material mixture during a process involving frictions can be suppressed, to improve yields.

[9025] For example, in the case of negative electrode, when a needle-shaped graphite is used for the active under nature and compared with the case when a fishe graphite is used, the surface roughness of the surface of the negative electrode increases, causing a tendency to decrease yields. In such a case as well, by decreasing the surface roughness of the protous film formed on the surface of the negative electrode, the separation of material mixture can be prevented effectively. The separation of material mixture is

likely to occur when a filler harder than the electrode active material is disposed along the asperities of the electrode surface.

[9026] By making the surface of porous film more smooth than the electrode surface, which is the base of the porous film, separation of the material mixture due to friction, which can happen in the conventional lithium ion secondary abstray having no porous film as well, can be suppressed, to greatly decrease defects of internal short circuit caused mainly by separation, and to increase yields.

[0027] Based on such, an electrode plate with a greater surface roughness can be applicable to a practical use. Therefore, a material which has been considered inapplicable for usage conventionally can be selected as an active material, improving versatility of a high performance lithium ion secondary batters.

[0028] In order to sufficiently adhere the porous film to a surface of an electrode, which is a base, it is desirable that the content of the inorganic filter included in the porous film is 99 wt % or below.

[9029] It is desirable that in the proross film, the inorganic filter comprises an inorganic oxide. It is desirable that the surface of the inorganic oxide is alkaline, and that the inorganic oxide has a BET specific surface area of 0.9 m<sup>2</sup>/s or orner. It is know that when a filler having an alkaline site on the surface thereof and a polymer having an acidic group are mixed, ecided group of the polymer makes bonding with the alkaline site of the filler (Ref. Development Technology of Functional Filler, CMC Publishing CO, LTD, pp. 37-47).

[0030] Many binders produce an acidic group by colations or reduction in a battery. Among the binders, nabber particles are expecially apt to hold an acidic group. When producing a battery having the activated binder group, in many cases, rubber particles are used as an electrode binder group, in many cases, rubber particles are used as an electrode binder, since the electrode plain each to have flexibility. When the acidic group captures lithium ion in the battery, desired buttery performance cannot be obstanced, and the life of nibber particles is shortened at the same time. On the other nabes brainly with the elidities is do of the tiles, the desired group is suppressed that to a formation of a block the acidic group is suppressed that to a formation of the shock group is suppressed that to a formation of the shock elicities.

[0031] It is desirable that the inorganic oxide includes at least one selected from the group consisting of alumina and titanium oxide. In view of obtaining sufficient electrolyteresistance, it is desirable that α-alumina is used particularly. among alumina. Also, since the particle size of titanium oxide can be easily controlled to 0.1 µm or below relatively, titanium oxide is suitable for controlling the filled in structure, porosity, and pore size of the porous film. Since titanium oxide has a lower degree of hardness compared with alumina, which is also used as abrasives, titanium oxide is also excellent with regard to the point that a kneader pot will not be damaged at the time of preparing a material paste for porous film. Although a usage of silica as an inorganic oxide has also been proposed, alumina and titanium oxide are preferable in view of battery life and reliability, since silica may be eroded by non-aqueous electrolyte.

[0032] It is desirable that the inorganic filler comprises a mixture of a large particle group and a small particle group, and the average particle size A of the large particle group and the average particle size B of the small particle group satisfy the formula (1):

0.05≤B(A≤0.25

## BRIEF DESCRIPTION OF DRAWINGS

[0033] FIG. 1 is a view schematically showing a structure of a porous film of the present invention.

[0034] FIG. 2 is a view schematically showing an example of electrode arrangement in which a porous film of the present invention is adhered to the electrode.

[0035] FIG. 3 is a vertical cross sectional view of an example of a lithium ion secondary battery of the present invention

[0036] FIG. 4 is a schematic view showing a structure of a conventional porous film.

[0037] FIG. 5 is a schematic view showing a structure of another conventional porous film.

[0038] FIG. 6 is an example of the FT-IR absorption spectrum of an example of a first rubber including an acrylonitrile unit (core-shell type particles).

[0039] FIG. 7 is an SEM photograph of a cross section of a negative electrode of the present invention.

[0040] FIG. 8 is an SEM photograph of a cross section of a porous film of the present invention.

# BEST MODE FOR CARRYING OUT THE

[0041] In the present invention, a porus film interposed between a positive electrode and a negative electrode comprises an inorganic filler and a first binder. The porous film has to be adhered to a surface of at least the negative electrode. This is because in lithium in secondary batteries, the width of the negative electrode is designed to be largethan, the width of the positive electrode, to present nelectric current from concentrating on an edge of the negative electrode.

[0042] Even though a porous film is formed on a sheet separator, the porous film shrinks when the separator shrinks. Therefore, when a great amount of heat is generated by a short circuit, the shrinkage of the porous film with the separator cannot be avoided, regardless of the heat resistance of the provus film itself.

[9043] Also, when an independent sheet composed of a sole porous film is to be formed, in view of keeping its sheet form, its thickness should be made very large, and a great amount of the binder becomes necessary. Therefore, in view of battery characteristics and design expactive, it is less practical to form an independent sheet of the porous film above.

[0044] Additionally, by forming the porous film adhered to a surface of an electrode, aspertites of the surface of an electrode aspert an extra electrode material mixture layer are covered with the porous film, making the friction of electrode surface small. As a result, a separation of the material mixture can be suppressed effectively.

[0045] The first binder content in the porous film needs to be 1.5 to 8 parts by weight per 100 parts by weight of the

filler. When the first binder content is less than 1.5 parts by weight, a portous film with sufficient strength cannot be obtained, and when the first binder content is over 8 parts by weight, gaps in the porous film become insufficient, decreasing lithium in permeability to lower rate characteristic. Also, a control of the microporous structure formed by the gaps among filler particles becomes difficult.

[0046] The first binder has to be water-insoluble.

[0047] The reason is that the most of high performance negative electrodes include a water-soluble polymer such as cellulose resin as a thickener. If the first hinder is water soluble, the first hinder has to be dissolved in water at the time of peparing a raw material paste for protus film. When time of peparing a raw material paste for protus film, when such raw material paste is applied on the negative electrode, the water-soluble polymer in the negative electrode weekly by water included in the raw material paste for protus film. In this case, the negative electrode deforms to cause a disadvantage such as a drawtie decrease in vields.

[0048] Herein, "the binder is water-insoluble" means that a substantially uniform solution cannot be obtained even the hinder is mixed with water. On the contrary, it is desirable that the binder dissolves homogenously in an organic soluted

[0049] In the porous film, it is prefemble that a robber (a filter tribber) including an arcylomitrile unit, desirably a polyacy-jointrile chain, is used as the first hinder entirely or partly. The first nubber including a polyacy-polyatric hain, is used as the first hinder entirely or partly. The first nubber including a polyacy-polyatric hain has a higher decomposition temperature, due to the heat resistance inhoratin in polyacy-polyatric (the softening point of 250 a 50° C., the decomposition temperature of 350° C.).

[9050] It is preferable that the first rubber is particular, When the first rubber is particular, poin-sablesoin some possible between filter particles, so that sufficient binding effect can be obtained even with a small amount thereof. The sufficient particles are of point-saflession is schematically shown in FIG. 1. Since filter particles 2 fare adhered to each other by a first bright particles 24 fare adhered to each other by a first bright FIG. 1. Since filter particles 24 fare adhered to each other by a first bright FIG. 1. Since filter particles 24 fare adhered to each other by a first bright FIG. 1. Since filter particles 24 fare adhered to each other by a first bright FIG. 1. Since filter particles 24 fare adhered to each other by first filter fil

[0051] It is desirable that the first rubber is core-shell type rubber particles having an adhesive surface portion. This is hecause the core-shell type rubber particles can exert sufficient binding effect even with a small amount thereof, securing more gaps in the portous film and sufficient moving path for the liquid electrolyte or lithium ion.

[0052] It is preferable that the core-shell type rubber particles further include an acrylate unit, other than the acrylating unit. Additionally, it is preferable that the acrylate unit forms the adhesive surface portion. For the acrylate unit, 2-ethylhexyl acrylate is suitable.

[0053] The binder having rubber elasticity is excellent also in the sense that impact resistance is given to the porous film. The porous film including the first rubber as a hinder is resistant to cracking when winding the positive electrode

and the negative electrode. Therefore, higher yields for a battery including a wound-type electrode plate group can be maintained. On the other hand, in the case of the porous film including a hard binder with no rubber elasticity, there is a concern for occurrence of cracks during the manufacturing process of a wound-type hattery.

[0054] As preferable examples of the first rubber, modified acrylonitrile rubbers such as BM-500B and BM-720H manufactured by Zeon Corporation can be mentioned, for example, and these are commercially available.

[0055] When using a first hinder comprising plural kinds of resin material for the protous film, it is preferable that her ratio of the first rubber relative to the first binder as a whole is 20 to 80 wt 56. When the first rubber is particulate, the average particle size of the particles is preferably 0.05 to 0.3 µm, in terms of obtaining a porous film with well-balanced strength and porosity.

[0056] When selecting the resin material used for a buttery, the stability of resin derived from Molecular One Motery of the most of the Molecular One of the Comtionally, Based on such indicator, generally, single comtonally, Based on such indicator, generally, single component resins or a combination thereof (copolymen) is selected. Therefore, nobes including a polysecy iontritic chain, which is unstable under the negative electrode potential, is hardly selected, from a conventional viewpoint.

[9057] When a first binder comprising plural kinds of reammental as used in the porous filing, of the reain material unter that the first rubber, fluorecarbon resis such as poly-winglishen fluoride (PVDF), collusion resis such as poly-winglishen fluoride (PVDF), collusion resis such as car-boxymethyl cellulose (CMC), or polyvinglyrmolindar (PVP) can be used. Also, in view of giving an appropriate viscosity to the num material paste for porous film, it is preferable that fluorocarbon resis (for example, PVDF) with a molecular weight of 100000 to 1000000) is used in combination with the first rubber.

[0058] In the core-shell type rubber particles including a polyacrylonitrile chain and an acrylate unit, in terms of balance hetween adhesiveness and rubher elasticity, it is preferable that the absorption intensity based on C=() stretching vibration is 3 to 50 times the absorption intensity based on C=N stretching vibration of the acrylonitrile unit, in absorption spectrum of rubber particles obtained by an FT-IR measurement. When the absorption intensity based on C=O stretching vibration is less than 3 times the absorption intensity based on C-N stretching vibration, binding effect of the rubber particles becomes insufficient, and when the absorption intensity based on C=O stretching vibration is over 50 times, rubber elasticity of the rubber particles hecomes insufficient, to weaken the strength of the porous film. The absorption intensity refers to the height of absorption peak seen from the base line of the spectrum.

[9059] In the FT-IR measurement, absorption spectrum of the core-shell type rubber particles can be measured by using a sample in which the rubber particles are applied on a RBr plate. for example, Generally, the absorption based on C=0 stretching vibration is observed around TD0 to 1760 cm<sup>-1</sup>, and the absorption based on C=N stretching vibration is observed around 2200 to 2280 cm<sup>-1</sup>.

[0060] In the porous film, the first rubber has to have a decomposition temperature of 250° C, or more. Also, when

the first rubber particles are crystalline, the first rubber has to have a crystalline melting point of 250° C, or more.

[9061] The reason is that in a nail penetration test, which is seven as an evaluation regarding an internal short circuit, the temperature at the time of an internal short circuit, the temperature at the time of an internal short circuit, the temperature and the conditions. In such a high temperature, the first rubber having a decomposition temperature below 250°C, or the first rubber having a crystalline melting point of helow 250°C. 6 bitmed out or excessively softened, to deform the porus film deformation of the portous film causes abnormal overheating.

[0062] It is desirable that the inorganic filler included in the porous film has a heat resistance equal to or higher than that of the first binder, is stable electrochemically under the environment of the lithium ion secondary battery in operation, and is suitable for making it into a neste (paint).

[0063] The BFT specific surface area of the filter has to be on m'gs or more, preferably 1.5 m'gs or more, in view of making the impregnation of electrolyte into the electrode plate group easier, while improving battery performance and file. When the BFT specific surface area is below 0.9 m'g, the adhesion between the first hinder and the filler denorates to weaken the strength of the porous film, and especially in manufacturing a wound-type electrode plate group, causing a deflective product. Additionally, it is preferable that the BFT specific surface area is not too large, and is 150 m'g or below, in view of suppressing congulation of the more sufficient of the procuss film of the plate of the production of the procuss film primaring fluidity of the raw material paste for more sufficient.

(9064) It is preferable that the specific gravity of the minorganic filter is  $0.3 \times 0.5$  g/cm<sup>2</sup> in view of shortenian incorpanic filter is  $0.3 \times 0.5$  g/cm<sup>2</sup> in view of shortenian kneading time of the raw muscrial pase for porous film. Further, it is preferable that the average particle size of preferable that the average particle size of preferably  $0.2 \times 0.1 \times 0.5$  m, and further perferably  $0.2 \times 0.1 \times 0.1 \times 0.1 \times 0.1$  m, and further perferably  $0.2 \times 0.1 \times 0.1 \times 0.1 \times 0.1$  m, and further perferably  $0.2 \times 0.1 \times 0.1 \times 0.1 \times 0.1$  m and further perferably  $0.2 \times 0.1 \times 0.1 \times 0.1 \times 0.1$  m in sidilicult to be formed. When waverage particle size is too small, the necessary amount of the resin anterial increases with the increase in the surface are of the filter, thereby making it difficult to form sufficient sums in the propure lim.

[9065] Also, in view of controlling the filled-in state of the filter, it is preferable that the filter comprises a mixture of a large particle group and a small particle group. It is preferable that the average particle size A (volume-based  $D_{\rm ej}$ ) of the large particle group is 0.2 to 2  $\mu$ m. Additionally, the average particle size B (volume-based  $D_{\rm ej}$ ) of the small particle group is 0.2 to 0.01 to 0.5  $\mu$ m.

[9066] It is preferable that the average particle size A of the large particle group and the average particle size B of the small particle group satisfy the formula (1):

#### 0.05 SB(A S0.25)

[9067]. When the value of B/A is helow 0.05. the surface area of the filler increases, which necessitates a great amount of the first binder in order to obtain a persons film with sufficient strength. Also, when the value of B/A is over 0.25, the gaps formed in the filler become excessively large, laiting to obtain capillary effect sufficiently, further decreasing the rate characteristic. [0068] It is preferable that the content of the small particle group in the filter as a whole is 1 to 20 tr 9 to, and that the rest is the large particle group. When the content of the small particle group is too small, it becomes difficult to fill the filter in the state close to the closest packing. When the content of the small particle group is too large, the surface area of the filter increases, necessiting a large amount of the first binder in order to obtain the porous film with sufficient strength.

[0009] For the above inorganic filler, inorganic oxides such as silicon oxide, alumina (Al,O<sub>2</sub>), and titamium oxide are preferable. Especially, shumina and ittamium oxide are preferable. Especially, shumina and ittamium oxide are preferable, and among titamium oxide, anattuse-type is preferable, rather than rutile-type. The surface of the anattuse-type titamium oxide has an alkaline site, which makes a bond with an acidic group included in the first binder to stabilize the structure of the porous film. With the use of annatse-type titamium oxide, a porous film with the use of annatse-type titamium oxide, a porous film with cellent shape-releation, adhesion, electrolyte-resistance, and potential cresistance are also commented. Here the control of the preference are also commented with the control of the control of the preference are also commented with the control of the control of the control of the control of the preference are also commented with the control of th

[0070] The inorganic oxide can be used alone, or can be used in combination of two or more. However, the content of the alumina or of the titanium oxide in the filler as a whole is preferably 50 wt % or more. A plurality of prorus films including different kinds of fillers may be laminated as well.

[0071] The following can be mentioned as the reasons why alumina is preferably used.

[0072] (1) The median diameter of alumina is suitable for forming the microporous structure (median diameter 0.02 to 0.09 µm) required for the porous film.

[0073] (2) Alumina is stable for both potentials of oxidation and reduction (0 to 5 V/vs 1.i).

[0074] (3) Alumina has less asperities in its particle surlace (has a small surface area), and a porous film with a higher strength can be easily obtained by a small amount of binder usage.

[9075] It is preferable that gaps in the porous film have the pore size of 0.1 μm (100 nm) or below, and further preferably 0.02 to 0.07 μm (20 to 70 nm), since when the pore size or distribution of the gaps in the porous film is not uniform, this meaning the gaps in the proper film is not uniform, the gaps are distributed uniformly. Also, it is desirable that such gaps are distributed uniformly.

[9076] In view of making the pore size or distribution of gaps in the porous film uniform, it is desirable that titanium oxide with relatively small particle size is used. To be specific, the particle size of itianium oxide is preferably 0.1 um or below. Also, it is preferable that the average particle size (median diameter) based on the number of titanium oxide is 0.04 to 0.1 um.

[0077] When ilmenite is used as a raw material of titanium oxide, ilmenite is dissolved by sulfuric acid in a reaction vessel, and then a step of sedimentation is carried out for separating the dissolved matter. The particle size of titanium oxide can be controlled by the reaction time at this time. [0078] Among the first binders, many are oxidized or reduced in a battery to produce an acide group. Particularly, rubber particles easily carry the acide group, la the lithium ion secondary battery, battery performance deteriorates when the acidic group captures lithium ion. Therefore, it is preferable that the acidic group in the first binder is decreased as much as possible, by bonding the acidic group of the first binder with the filter surface.

[0079] On the other hand, some filters are alkaline on a surface thereof. When the filter has an alkaline site on the surface thereof, the bonding strongth between the acidic group of the first binder and the surface of the filter is begittened. The alkaline site is formed from cations such as aluminum ion. The filter surface preferably has a pH of over 7 and not larger than 12.

[0880] The porous film can be obtained in a state such that it is athered to an electrede surface, by a step of applying a paste including the raw material of the porous film (hereinar referred to as raw material pasts) on the electrede surface. In the lithium ion secondary battery, the width of the negative electrode must be designed to become wider than the width of the positive observable, in order to prevent a negative electrode. In order to prevent a negative electrode. Therefore, in terms of fill-self-of-the order of the negative electrode. Therefore, in terms of fill-self-of-the negative electrode chaving a larger width.

[0081] The raw material paste for porous film is prepared by dispersing the filter and the first binder in a liquid component. For the liquid component, N-methyl-2-pyrotidone, acetone, or lower alcohol can be used, and nonaqueous electrolyte can be used as well.

[0082] The content of the raw material (a total of the filler and the first binder) in the raw material paste for the porous film is preferably 25 to 70 wt %. When the raw material content is too small, the porous film with a desired thickness and strength is difficult to be obtained, and when the raw material content is too large, the viscosity of the paste becomes higher to make the application difficult.

[9083] Although the thickness of the porus film is not particularly limited, the thickness is perferably 0.5 to 20 µm, in view of making full use of the safety improvement function by the porous film, while maintaining the designed capacity of the battery. Also, when a sheet separator generally used currently is used in combination with the portus film, the total thickness of the sheet separator and the portus film can be controlled to become 15 to 30 µm.

[0084] The usage of titanium oxide having a small particle size for the filler is advantageous for making the porous film than, since the pore size of the gaps in the porous film can be controlled to become small. That is, by using titanium oxide having the small particle size the spatial share of electrodes in a battery-can may be larger to increase the capacity, since a certain level of performance can be obtained even when the thickness of the porous film is thinner.

[0085] The positive electrode and the negative electrode capable of absorbing and desorbing lithium ion usually comprise an electrode contains a material and an electrode material mixture raried thereon. The electrode material mixture includes at feast an active material and a binder, and includes a conductive agent as necessary. The electrode plate is

generally prepared by applying a mixture of the electrode material mixture and a dispersion medium (material mixture paste) to the electrode core material, drying the mixture, and then rolling.

[9086] In the present invention, by controlling the surface roughness of the proton film a higher production yield can be achieved. In particular, by controlling the surface roughness of the proton film to become smaller than an surface roughness of the electrode surface where the most surface andered, the yield can be improved. This is because the protons film causes a descrease of the friction on the material mixture layer surface, the friction causing the separation of the material mixture. In view of avoiding the separation of the material mixture effectively, it is desirable that the average surface roughness of the protons film Ra measurement instrument is below 2.1 µm, and further desirably below 1.5 µm.

[0087] For the particular method to form, on the material mixture layer, the protos film having a smaller surface roughness than that of the material mixture layer, methods of applying the my material pasts for porous film on the material mixture layers such as dic coating method or gravare printing are easy. Also, a method of immersing the electrode plate in the raw material pasts such as a batch style or successive style dip method, can be adopted. Further, a method of spraying the raw material pasts on the material mixture layer such as spray emitting method can also be adopted. In any of these methods, an optimal viscosity of the row material pasts should be determined from the relationship between the target thickness of the porous film and the apprities of the surface of the electrode plate as the base.

[0088] A cross-sectional photo by scanning electron microscope (SEM) of an example of a negative electrode plate of the present invention is shown in FIG. 7. The upper part and the bottom part of FIG. 7 are almost symmetrical. and an area in the center shown by a near-white color is a negative electrode core material. The areas shown by a near-black color on both sides of the negative electrode core material are the negative electrode material mixture layers, respectively. On the outside of each negative electrode material mixture layer, porous films are formed. Additionally, a cross-sectional photo by SEM including an interface between the porous film and the negative electrode material mixture layer is shown in FIG. 8. In the negative electrode material mixture layer as the base (the bottom part of the FIG.), only a flake graphite is used as an active material. From these FIG.s, it is clear that by optimizing the viscosity of the raw material paste for porous film, the raw material paste can enter into minute asperities of the negative electrode to smooth out the electrode surface.

[0089] Generally, a negative electrode includes a negative electrode active material, a binder (a second binder), and a water-soluble polymer.

[0090] For the negative electrode active material, various natural graphites, various artificial graphites, silicon-containing composite materials such as silicide, and various alloy materials can be used.

[0091] When controlling the surface roughness of the porous film, a vapor grown carbon fiber (VGCF) which is especially highly conductive among negative electrode

active materials can be used in a larger amount than conventionally. This is because although the VGCF causes many aspertites on the material mixture layer surface, such aspertites are to be covered by the porous film to avoid the problems such as the separation of the material mixture.

[9092] A short circuit due to deposits of metallic lithium in the porous film appears to occur mainly when lithium acceptability of the negative electrode is inferior to lithium permeability of the porous film. Lithium acceptability of negative electrode decreases as the amount of the second binder increases, to raise possibility of the short circuit occurrence.

[9093] Therefore, in the present invention, it is desirable that the decrease of lithium ion acceptability of negative detected is suppressed to the minimum by using the second binder which can exert sufficient binding effect with a small amount of Isage, in order for that, it is effective to use rubber particles (second rubber particles) in combination with a water-soluble polymer as a second binder.

[0094] For the second rubber particles, rubber particles including a styrene unit and a butadiene unit are preferable. For example, styrene-butadiene copolymer (SBR), and modified SBR can be used, although not limited thereto.

[0095] For the water-soluble polymer, cellulosc resins are preferable, and those including a methylcellulose unit is particularly preferable. For example, carboxymethyl cellulose (CMC), methylcellulose, and metal salts thereof are used preferably. Among these, CMC partly converted into alkali metal salts is the most preferable.

[0096] The amount of the second binder and the amount of the water-soluble polymer included in the negative electrode are preferably 0.1 to 5 parts by weight and 0.1 to 5 parts by weight, respectively, per 100 parts by weight of the negative electrode active material.

[0997] In order to keep fithium acceptability of the negative electrode at a higher kevel, it is desirable that the total amount of the second rubber particles and the water-solids polymer included in the negative electrode is 1.5 to 3 bars by weight per 100 parts by weight of the negative electrode active material. When the total amount of these is below 1.5 parts by weight per 100 parts by weight of the negative electrode active material. Gestraction of the structure of the negative electrode may occur (material mixture peelings), and when over 3 parts by weight of the negative electrode active material. Infulum acceptability of the negative electrode active material. Infulum acceptability of the negative electrode active material. Infulum acceptability of the negative electrode active material.

[0098] It is preferable that the average particle size of the second rubber particles is 0.1 to 1 µm, in view of securing binding effect, while avoiding excessive coverage of the active material thereby.

[0999] It is preferable that weight ratio of the second intheir particles and the water-soluble polymer is 1:10 to 10.1. When the ratio of the water-soluble polymer relative to the rubber particles is too large, then to poor flexibility of the water-soluble polymer, the flexibility of the electrode plate the water soluble polymer, the flexibility of the electrode plate deteriorates, making the material mixture prone to peel atteted turne of forming the electrode plate group. On the other hand, when the ratio of the second rubber purifice relative to the water-soluble polymer is too large, stability of the negative electrode material mixture paste prepared in the production process of the electrode plate declines, to cause variations in the applied amounts of the paste on the electrode core material, and to decrease productivity.

[0100] Generally, the positive electrode includes at least a positive electrode active material, a positive electrode binder, and a conductive agent.

[9101] For the positive electrode active material, a composite oxide can be mentioned. For the composite oxide, lithium cobaltute, modified lithium cobaltute, lithium nickelate, modified lithium nickelate, lithium manganate, modified lithium manganate are preferable. Each modified product may include elements such as aluminum, magnesium, or the like. Also, some include at least two selected from cobalt nickel, and manganese.

[9102] For the positive electrode binder, atthrough not particularly limited, polyterfullasoverlybene (PTFP), modified excylonitrile rubber particles (BM-500B manufactured by Zona Corporation), and polyvinytidene fluorde (PV1FF) can be used. It is preferable that PTFH and BM-500B are used in combination with CMC, polytythyene oxide (PEO), or modified acrylonitrile rubber (BM-720H manufactured by Zona Corporation), which serves as a theckner for the armount of the properties of the properties

[0103] For the conductive agent, acetylene black, Ketjen black, and various graphites can be used. These can be used alone, or used in combination of two or more.

[0104] A non-aquous electrilyte generally comprises a littims and and anon-aquous solven. For the littims and in littim and an anon-aquous solven. For the littim and in littim and and solven, delyten exchange (FC, LiBF, is used, for example, For the non-aqueus) solven, delyten exchange (FC, dinethyl carbonate (DEC, dine

[0163] Vinylene carbonate (VC), cyclohexylbenzene (CHB), and modified VC or CHB can be used, in order to form an excellent film on the positive electrode and/or the negative electrode for securing stability at the time of overcharge.

[016] A positive electrode capable of absorbing and desorbing lithium ion, negative electrode capable of absorbing and desorbing lithium ion, and non-aqueous electrody can be appropriately selected from known materials for usage, and are not limited to the above listed ones.

[0107] In the present invention, by using a procus film excellent in heat resistance and a conventional sheet separator, safety can be achieved at a higher level. Although the sheet separator is not particularly limited as long as the separator comprises a material endurable to the usage enrish connents of the lithium into battery, generally, nicreportous film comprising an otefan resin such as polyethylene or polyprepylene is used. The microprorus film can be a single layer film comprising one kind of olefin resin, or can be a plural layer film comprising one find of olefin resin, or can be a plural layer film comprising to more ofetin resin as

[0108] Although the thickness of the sheet separator is not particularly limited, in view of maintaining the designed capacity of the battery, the thickness is preferably 8 to 30 am.

[0109] In the following, the present invention is concretely described based on Examples.

[0110] First, the method of physical property evaluation for the filler and the first binder forming the porous film used in Examples is described.

#### [1] pH of Filler

[0111] For p11 measurement, a boiling extraction glass electrode measurement method was used, and the measurement was carried out based on JIS R6129-1976 and JIS28802-1984. Also, pH measurement using ELS-8000 (pH titration apparatus) manufactured by Otsuka Electronics Co., Ltd. was carried out.

[2] BET Specific Surface Area of Filler

[0112] The measurement of the BET specific surface area was carried out by using a direct-reading specific surface area measurement apparatus, based on BET single point method. First, 0.5 to 1 g of insulating filler as a sample was placed in a glass cell, and cleaning was carried out for 20 to 30 minutes at 250° C, under a flow of a mixed carrier gas of nitrogen and helium (volume ratio N3:He=30:70). Then, while cooling the sample of the insulating filler with liquid nitrogen, N2 in the carrier gas was adsorbed thereon. Subsequently, the temperature of the sample of the insulating filler was raised to give room temperature. The desorbed amount of N, was detected by a thermal conductive detector, and the specific surface area was calculated from the surface area corresponding to the desorbed amount of N2 and a mass of the sample after the measurement. For the calculation, NOVA2000 manufactured by Yuasa Ionics Inc. was used.

#### [3] Heat Resistance of Filler

[0113] Differential scanning calorimetry (DSC) and thermogravimetry-differential thermal analysis (TG-DIA) were carried out for the sample of the filler, to evaluate heat resistance based on the temperature of an inflection point in the DSC measurement or the temperature at the start point of the weight change in the TG-DTA measurement.

[4] Crystalline Melting Point or Decomposition Temperature of Binder

[0114] Differential scanning calorimetry (DSC) and thermagnavimetry-differential thermal analysis (TcD-DTA) of the hinder sample were carried out, and the temperature of the inflection point in the DSC measurement or the temperature at the start point of the weight change in the 10-311 measurement was regarded as the crystalline melting point or the decomposition temperature.

[5] Affinity between First Binder and Water

[0115] Solubility of the first binder in water at room temperature and atmospheric pressure was measured, and it was determined as "water-insoluble" when the solubility is I wt % or below.

[6] Particle Size Distribution and Median Diameter of Filler

[0116] The measurement of the average particle size was carried out by using a laser diffraction scattering particle size distribution analyzer (Master Sizet 2000 manufactured by Malvern Instruments Ltd.) and a centrifugal particle size distribution analyzer (SA-CP3 manufactured by Shimadzu Corporation).

## EXAMPLE 1

[0117] FIG. 2 and FIG. 3 are referred to in the following explanation.

(a) Preparation of Raw Material Paste of Porous Film

[0118] A raw material paste for a perous film was prepared by dispersing a filter and a first binder in N-methyl-2pyrrolidone (NMP) in the ratios shown in Table 1, and then kneading them. The total content of the filter and the first binder in the paste was set as 50 mt % in any of the case.

[0119] For the first binder, core-shell type rubber particles including an acrylonitrile unit (a first rubber) and polyvinytidene fluoride (PVDF) with a molecular weight of 350, 000 were used in combination.

[0120] Herein, for the core-shell type rubber particles, BM500B (average particle size 0.2 µm) manufactured by Zeon Corporation, comprising aerylonitrile-aerylate copolymer was used.

[0121] Physical properties of BM500B are shown in the following.

[0122] <1> Crystalline Melting Point: None (amorphous)

[0123] <2> Decomposition Temperature: 308.5° C.

[0124] <3> Affinity With Water: Water-Insoluble

[0125] The absorption spectrum of the first rubber (BM500B) obtained by an FF-IR measurement is shown in FIG. 6. For the measurement apparatus, FF-IR microscope (Continuum manufactured by Nicolet Instrument Corporation, Bight source: AWATAR-360) was used.

[0126] The measurement conditions were set as: sample sean 32, background scan 32, resolution 4000, and sample gain 1.0. For the measurement sample, the first rubber was dispersed in N-methyl-2-pyrrolidon, applied on a KBr plate, and dried for the usage.

[9127] In FIG. 6, the absorption peak observed near 2240 cm<sup>-1</sup> is based on C≥N stretching vibration of acrylonitrile, and absorption peak near 1733 cm<sup>-2</sup> is based on C≡O stretching vibration. In FIG. 6, the absorption peak intensity (peak height) based on C=O stretching vibration is about 10 times the absorption peak intensity (peak height) based on C=O stretching vibration is about 10 times the absorption peak intensity (peak height) based on C=N stretching vibration of acrylonitrile unit.

[9128] For the filter, Al<sub>2</sub>O<sub>2</sub> was used, Herefa, "alumina a" with the average particle size of 0.2 to μm was used to 2 μm used with the average particle size of 0.0 to μm was used to 0.4 μm and "alumina b" with the average particle size of 0.0 to 0.15 μm was used. The "alumina b" content and the off 0.0 to 0.15 μm was used. The "alumina b" content and the manual bus was used to 0.0 to 0.5 μm was used. The "alumina b" content and the manual bus was used to 0.0 to 0

(b) Fabrication of Positive Electrode

[0129] To 100 parts by weight of LiCoO<sub>2</sub>, 4 parts by weight of polyvinylidene fluoride (PVDF) as a binder and 3

parts by weight of acetylene black as a conductive agent were added and subsequently an appropriate amount of N-methyl-2-pyrodistone (NMP) was added and then cancelled, to prepare a positive electrode material mixture paste frace obtained positive electrode material mixture paste. The chained positive electrode material mixture paste 21 with a histonicos of 20 µm, and 21 with a histonicos of 20 µm, and 22 when a histonicos of 20 µm, and chansity of the active material (density of LiCoO<sub>2</sub>) in a positive electrode anterial mixture 22 became 33 µm, to produce a positive electrode 23. A positive electrode lead 24 made of aluminum was connected to the positive electrode lead

#### (c) Fabrication of Negative Electrode

[0130] To 100 parts by weight of spherical artificial graphice, as a second binder, rubber particles comprising a copyolymer including a styrene unit and a batadiene unit, and carboxymethyl cellulose (CMC) were added in the ratios shown in Table 1. An appropriate amount of water was added thereto and then kneaded, to prepare a negative electrude material mixture paste.

[0131] Herein, for the rubber particles comprising a copolymer including a styrene unit and a butatienc unit, BM4003 (average particle size 0.4 µm) manufactured by ZEON Corporation, comprising a styrene-methacrylic acid-butadiene copolymer, was used.

[0132] The obtained negative electrode material mixture pasts was applied onto one side of a copper foil core material. 25 with a thickness of 15 µm, and then rolled so that the density of the active material (density of graphite) in the negative electrode material mixture 26 became 1.4 g/ml, to produce a negative electrode. To the negative electrode 27, a negative electrode lead 28 mode of cupper was connected.

## (d) Formation of Porous Film

[0133] Then, the raw material paste for porous film was applied to give a thickness of 20 µm on one side of the negative electrode 27 so that the negative electrode material mixture 26 was completely covered therewith, to form a porous film 31.

## (e) Preparation of Electrolyte

[0134] For the non-aqueous electrolyte, to a mixed solvent of ethylene carbonate, ethylmethyl carbonate, and dimethyl carbonate with a volume ratio of 11:11, lithium hexafluor-phosphate (LiPF<sub>6</sub>) was dissolved to give a concentration of 1 mol/L. Also, to the mixed solvent, 4% by volume of vinvlene carbonate was added.

#### (f) Fabrication of Battery

[0135] As shown in FIG. 2, the positive electrode 23 was disposed on the porous film 31 to form a stacked-type unit cell comprising a pair of positive electrode and negative electrode. This unit cell was enveloped by an outer jacket 32 comprising aluminum laminate sheet, and then, a nonangueous electrotyle was charated into the outer jacket.

[0136] Then, resin sealing materials 33 overing a part of the positive electrode lead 24 and the negative electrode lead 25 are dispared to be located at an opening end of the outer lacket 32, respectively, and the outer jacket 32 was senter lacket 32, respectively, and the outer jacket 32 was senter under vacuum while free ends of the respective lead were drawn to the outside. A lithium is necendary battery, the above the control of the control of the second of the second that the control of the second position of the second position.

TABLE 1

	Amoun Secon Binden/ parts weight Activ Mater (parts weigh	nd 100 by of re ind by	Amou Fin Binder parts weigh Fill (parts weigh	st /100 by it of or by	Avera	ge Particle Size of Filler (um)	Peeling of Negative	Peeling of Porous	Short	High- rate Char- ac- teris- tic
Example	BM400B	CMC	BM500B	PVDF	Alumina a	Alumina b	Electrode	Film	Circuit	(%)
X1	1	1	2	2	0.4		Absent	Absent	Absent	86.3
X2	0.75	0.75	2		0.4		Absent	Absent	Absent	88.1
Y1	0.5	0.5	2	2 2 2	0.4		Present	71000111	, tooth	00.1
X3	1.5	1.5	2	2	0.4		Absent	Absent	Absent	84.9
Y2	2	2	2	2	0.4		Absent	Absent	Present	
X4	1	1	0.75	0.75	0.4		Absent	Absent	Absent	89.9
Y3	1	1	0.5	0.5	0.4		Absent	Present		
X5	1	1	3	3	0.4		Absent	Absent	Absent	84.1
X6	1	1	4	4	0.4		Absent	Absent	Absent	80.5
Y4	1	1	5	5	0.4		Absent	Absent	Absent	73.8
X7	1	1	2	2	0.4	0.05(B/A = 0.125)	Absent	Absent	Absent	87.5
X8	1	1	2	2	0.4	0.02(B/A = 0.05)	Absent	Absent	Absent	89.1
X9	1	1	2	2	0.4	0.01(B'A = 0.025)	Absent	Slightly	Absent	89.3
X10	1	1	2	2	0.4	0.1(B/A = 0.25)	Absent	Absent	Absent	85.9
X11	1	1	2	2	0.4	0.15(B/A = 0.375)	Absent	Absem	Absent	84.8
X12	1	1	0.8	3.2	0.4		Absent	Slightly	Absent	86.2
X13	1	1	3.2	0.8	0.4		Absent	Absent	Absent	86.8
X14	1	1	2	2	0.2		Absent	Slightly	Absent	88.1
X15	1	1	2	2	1		Absent	Absent	Absent	86.2
X16	1	1	2	2	2		Absent	Absent	Absent	85.7

TABLE 1-continued

Example	Mater (parts weig) BM400B	by	Weigh Fill (purti Weigh BM500B	er by		Particle Size of Filler (turn) Alumina b	Peeling of Negative Electrode	Peeling of Porous Film	Short Circuit	Char- ac- teris- tic (%)
	Secon Binder parts weight Activ Mater	100 by r of re	Amou Fir Binder parts weigh Fill	st r100 by it of	Average	: Particle Size	Peclino	Peeling		High- rate Char-

#### (Evaluation)

[0137] Evaluations were carried out for the porous film prepared in the above and the completed lithium ion secondary battery, as in the following.

## [Presence of Absence of Peelings]

[0138] Appearances of the negative electrode and the porous film were observed to check if peelings occurred.

#### [Occurrence of Short Circuit]

[0139] Each hattery was charged at 120 mA till the battery voltage reached 4.2 V, and the charged battery was left at the atmosphere of 45° C. for a week. The battery voltage was measured afterwards, and it was determined that a short circuit occurred when the voltage was below 4.0 V. The results are shown in Table 1.

#### [High-Rate Characteristic]

[0140] After activating each hattery by carrying out a predetermined chape and discharge the batteries were charged at 120 m. Ait II the hattery voltage reached 4.2 V, and then discharged at 60 m. Ait III the battery voltage reached 4.3 V. Then, the same charging was varied out and discharging was carried out at 600 m.A III the battery voltage reached 3. V. Then, the ratio of discharge capacity when discharging at 60 m.A relative to the discharge capacity when discharging at 60 m.A relative to the discharge capacity when discharging in 180 m. S. The second of the capacity when discharging at 60 m. Awas obtained hypercentage. The results are shown in Table 1.

## (Results Consideration)

[0141] The results of Table 1 shows that the porous lilm with sufficient strength cannot he obtained when the amount of the first binder in the porous lilm is small, due to the occurrence of peclings of porous lilm. Also, it is shown that high-rate characteristic greatly declines when the amount of the first binder is too large. That is, the results of Table 1 imply that the content of the first binder in the porous film should be 1.5 to 8 parts by weight per 100 parts by weight of the filler.

[0142] On the other hand, it is shown that the negative observable with sufficient strength cannot be obtained when the amount of the second binder in the negative electrode is small due to the occurrence of the peelings of the negative electrode material mixture. Also, when the amount of the second binder is too large, a short circuit tends to occur due to the decline in lithium acceptability of the negative eletrode. When the battery in which it short circuit coccurwas disassembled to observe the cross section of the porous film, deposits of metallic lithium were seen in the porous lilm. Therefore, it can be seen that the amount of the second binder in the negative electrode should be 1.5 to 3 parts by weight per 100 parts by weight of the negative electrode active material.

[0143] It can be also seen that high-rate characteristic tends to gradually decline as the ratio (value B/A) of the average particle size of "alumina h" and "alumina a" increases. On the other hand, it can be seen that when the value B/A is too small, the strength of the porous film tends to decrease.

[0144] When the average particle size of the filter is too small, the surface area thereof becomes too large to cause insufficient first binder, showing a tendency to cause peelings of the porous film. On the other hand, when the filter is too large, the first binder becomes a surplus, showing a decline in high-rate characteristic.

#### EXAMPLE 2

## Battery A1

## (a) Preparation of Raw Material Paste of Porous Film

[0145] A raw material paste for prorus film was prepared by kneading 970 g of alumina with a median diameter of 0.3 µm as a filter, 375 g of BM-720H (MMP solution containing 8 wt % of a first rubber including a polyacrylonitrile chain as a first hinder) manufactured by JZBO Corporation, and an appropriate amount of NMP in a double-arm kneader.

[0146] Physical properties of BM-720H are shown below.

[0147] <1> Crystalline Melting Point: None (amorphous)

[0148] <2> Decomposition Temperature: 320° C.

[0149] <3> Affinity With Water: Water-Insoluble

#### (b) Fabrication of Positive Electrode

[0.150] A positive electrode material mixture paste was prepared by mixing 3 kg of lithium cohaltate, 1 kg of PVDP#1320 (MMP solution containing 12 wt % of PVDP) manufactured by Kurcha Chemical Industry Co., 1,td. as a binder, 90 g of seetly-lene black, and an appropriate amount of NMP, with a double-surn kneader. This paste was applied on an aluminum foil with a thickness of 15 µm, and rolled after drying, to form a positive electrode material mixture hyer. At this time, the thickness of the electrode plate comprising the aluminum foil and the material mixture layers was set as 160 µm. Afterwards, the electrode plate was slit to give a width which could be inserted into a can battery case of a cylindrical battery (product number 18650) to obtain a positive electrode horo.

#### (e) l'abrication of Negative Electrode

[0151] A negative electrode material mixture paste was prepared by mixing 3 kg of artificial graphite, 75 g of BM-400B (aqueous dispersion including 400 wt/8 of styleng-busidene copolymer (second rubber particle) as a second binder) manufactured by ZEON Corporation, 30 g of CMC as a water-soluble polymer, and an appropriate amount of water with a double-arm kneeder. This paste was applied on a copper foil with a thickness of 10 mm, and rolled after drying, to form a negative electrode material mixture layers was ext as 180 at this time, the thickness of the electrode plate comprising the copper foil and the material mixture layers was set as 180 which could be to electrode plate using the copper foil mixture layers was ext as 180 which could be to electrode plate using to gave a with which could be to electrode plate to mixture layers was extended by the control of the properties of the control of the mass for the control of the reason of the control of the mixture layers was extended to the control of the reason of the comprising the copper foil made to the control of the reason of the reason of the control of the reason of the control of the reason of the reason

#### (d) Formation of Porous Film

[0152] The raw material paste for porous film was applied on both sides of the positive electrode hoop, and then dried, to form a porous film having a thickness of 5  $\mu$ m per side adhered to the positive electrode surface.

#### (e) Preparation of Electrolyte

[0153] For the non-squeous electrolyte, a solvent mixture in which ethylene carbonate, ethyl methyl carbonate, and dimethyl carbonate were mixed with a volume ratio of 1:1:1, dissolving, lithium hexafluorophosphate (LiPF<sub>d</sub>) to give a concentration of 1 mol/liter was used. Also, 30% youlume of vinylene carbonate relative to the solvent mixture was added to the non-squeous electrolyte.

#### (f) Fabrication of Battery

[0154] The above positive electrode and negative electrode were cut to give a predetermined length, respectively, and were wound interposing a sheet separator with a thickness of 20 µm comprising a microperous film made of polyethylene, and then were inserted into the battery case. Then, 3.5 g of the showe electrolyte was weighted and injected into the battery case, and the opening of the case was seated. A cylindrical-shape 18650 thitum ion secondary battery Al was flus made.

## Battery B1

[0155] A battery B1 was made in the same manner as the battery A1, except that the porous film was not formed on both sides of the positive electrode hoop.

#### Battery B2

[0156]. The now material paste for porous film was applied on both sides of the sheet separator comprising a polyeth-ykene-made microporous film, and then dried, to form a porous film having a thickness of 5 µm per side adhered to the separator surface. A battery 182 was made in the same manner as the battery B1, except that the separator laving the porous film thus obtained was used.

#### Battery B3

[0157] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a prouse film having a thickness of 03 jum per side adhered to the negative electrode surface. A battery 133 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

#### Battery A2

[0158] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dired, to form a prous film having at thickness of 0.5 µm per side adhered to the negative electrode surface. A buttery A2 was made in the same manner as the buttery B1, except that the negative electrode having the porous film thus obtained was used.

#### Battery A3

[0159] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a prorous film having a thickness of 1 µm per side adhered to the negative electrode surface. A buttery A3 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

#### Battery A4

[0160] The raw material paste for porous film was applied no both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 5 µm per side adhered to the negative electrode surface. A battery A4 was made in the sume manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

## Battery A5

[9161] The raw material passe for porous film was applied to both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 10 µm per side adhered to the negative electrode surface. A battery A5 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

#### Battery A6

[0162] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 15 µm per side adhered to the negative electrode surface. A battery A6 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

## Battery A7

[0163] The raw material paste for porous film was upplied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 20 µm per side adhered to the negative electrode surface. A battery A7 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was next.

Battery B4

[9164] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried. to form a prouse film having a thickness of 30 µm per side adhered to the negative electrode surface. A battery B4 was made in the same manner as the battery B1, except that the negative electrode having the porous film thus obtained was used.

Battery B5

[0163] A battery B5 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 6 µm.

Battery A8

[0166] A battery A8 in which a thickness of the porous little adhered to the negative electrical surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 8 µm

Battery A9

[0167] A battery A9 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 10 µm.

Battery A10

[0168] A battery A10 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 15 µm.

Battery A11

[0169] A battery A11 in which a thickness of the porous film adhered to the negative electrods surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 25 µm.

Battery A12

[0170] A battery A12 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 30 µm.

Battery B6

[0171] A battery B6 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the thickness of the sheet separator was set as 40 um.

Battery B7

[0172] A battery B7 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 30 wt %.

Battery A13

[0173] A battery A13 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 50 wt %.

Battery A14

[0174] A battery A14 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 70 µm %

Battery A15

[0175] A battery A15 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 90 µm 9.

Battery A16

[0176] A battery A16 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 95 wt 96.

Battery A17

[0177] A battery A17 in which a thickness of the porous film adhered to the negative electrode surface was 5  $\mu$ M per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 99 wt %.

Battery B8

[0178] A battery B8 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that the filler content relative to the total of the filler and the first binder was set as 99.5 wt 9.

Battery B9

[0179] A battery B9 in which a thickness of the porsus thin adherent to the negative chectronic surface was find the filter of the negative chectronic surface was find the side thereof was made in the same manner as the battery Ascept that as the first binder, water-soluble CMC was sade instead of BM-720H to prepare the raw material passe for provise film. Since CMC was used as the first binder, in the battery B9, for the dispersion medium of the raw material passee, water was used instead of NMC CMC does not have a crystalline melting point, is amorphous, and has a decomposition temperature of 245° C.

Battery B10

[0.180] A battery B10 in which a thickness of the portus iffin adhered to the negative electroned surface was 5 miles film adhered to the negative electroned surface was 5 miles and except that as a first binder, water-instoable PVDF was except that as a first binder, water-instoable PVDF was except that as a first binder, water-instoable PVDF was consistent of instead of BM-720H, to prepare the raw material passe for procus film. PVDF has a crystalline melting pout and a decomposition temperature of 174° C, and 360° C, respectively.

## Battery B11

[0181] A battery B11 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery A4, except that as the filler, polyethylene beads with a median diameter of 0.3 µm were used instead of the alumina with a median diameter of 0.3 µm.

#### Battery A18

[0182] A battery A18 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side theroof was made in the same manner as the battery A4, except that as the filler, titania with a median diameter of 0.3 µm was used instead of the alumina with a median diameter of 0.3 µm.

## Battery A19

[9183] Upon making a negative electrode, instead of BM400F and CMC as the second binder, 8 wt \$50 (PC) relative to the artificial graphite was used. Herein, for PCDF, PVDF/130 with was used as the binder of the popular electrode was used. Except for the above, a battery, A19 in which a thickness of the prones film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery. A4

[0184] The configuration of the porous film is summarized and shown in Table 2. Additionally, the thickness of the sheet separator and the kinds of the second binder used for the negative electrode are summarized and shown in Table

TABLE 2

				Porous 1	ilm		
						Binder	
		Film	Fit	ler		Crystalline Melting	Decomposition
Battery No.	Adhered Portion		Kind	Content (wt %)	Kind	Point (° C.)	Temperature (° C.)
Al	Positive Electrode	5	Ahmuna	97	BM720H	None	320
42	Negative Electrode	0.5	Alumina	97	BM720H	None	320
A3	Negative Electrode	1	Alumna	97	BM720H	None	120
Α4	Negative Electrode	5	Alumina	97	BM720H	None	320
A5	Negative Electrode	10	Alumina	97	BM720H	None	320
A6	Negative Electrode	15	Alumins	97	BM720H	None	320
A7	Negative Electrode	20	Alumina	97	BM720H	None	320
A8	Negative Electrode	5	Alumina	97	BM720H	None	320
A9	Negative Electrode	5	Alumina	97	BM720H	None	320
A10	Negative Electrode	5	Alumina	97	BM720H	None	320
A11	Negative Electrode	5	Alumina	97	BM720H	None	320
A12	Negative Electrode	5	Alumina	97	BM720H	None	3.20
A13	Negative Electrode	5	Alumna	50	BM720H	None	320
A14	Negative Flectrode	5	Alumina	70	BM720H	None	320
A15	Negative Electrode	5	Alumina	90	BM720H	None	320
A16	Negative Electrode	5	Alumina	95	BM720H	None	320
A17	Negative Electrode	5	Alumina	99	BM720H	None	320
A18	Negstive Electrode	5	Titania	97	BM720H	None	320
A19	Negative Electrode	5	Alumina	97	BM720H	None	320
B1	None						
B2	Separator	5	Alumina	9?	BM720H	Nosc	320
B3	Negative Electrode	0.3	Alumina	97	BM720H	None	320
B4	Negative Electrode	30	Alumna	97	BM720H	None	360
B5	Negative Electrode	5	Alumina	97	BM720H	None	320

TABLE 2-continued

				Porous I	Film		
						Binder	
Battery No.	Adhered Portion	Film	Filler			Crystalline Melting	Decomposition
		Thickness (µm)	Kind	Content (wt %)	Kind	Point (° C.)	Temperature (° C.)
В6	Negative Electrode	5	Alumina	97	BM720H	None	320
В7	Negative Electrode	5	Alumina	30	BM720H	None	320
B8	Negative Electrode	5	Alumina	99.5	BM720H	None	320
B9	Negative Electrode	5	Alumma	97	CMC	None	245
B10	Negative Electrode	5	Alumna	97	PVDF	174	360
B11	Negative Electrode	5	PE beads	97	BM720H	None	320

[0185]

TABLE 3

			Porc	us Film		
Bat- tery No.	Separator Thickness (µm)	Negative Electrode Binder	Ad- hesive- ness	Flexibility (No. of Defects)	Negative Electrode Appearance	
Al	20	BM400B + CMC	OK	0	No Change	
A2	20	BM400B + CMC	OK	0	No Change	
A3	20	BM400B + CMC	OK	0	No Change	
Λ4	20	BM400B + CMC	OK	0	No Change	
Α5	20	BM400B + CMC	OK	0	No Change	
A6	20	BM400B + CMC	OK	0	No Change	
A7	20	BM400B + CMC	OK	0	No Change	
Λ8	8	BM400B + CMC	OK	0	No Change	
A9	10	BM400B + CMC	OK	0	No Change	
A10	15	BM400B + CMC	OK	0	No Change	
AH	25	BM400B + CMC	OK	0	No Change	
A12	30	BM400B + CMC	OK	0	No Change	
A13	20	BM400B + CMC	OK	0	No Change	
A14	20	BM400B + CMC	OK	0	No Change	
A15	20	BM400B + CMC	OK	0	No Change	
A16	20	BM400B + CMC	OK	0	No Change	
A17	20	BM400B + CMC	OK	1	No Change	
A18	20	BM400B + CMC	OK	0	No Change	
4.19	20	PVDF	OK	2	No Change	
B1	20	BM400B + CMC			No Change	
132	20	BM400B + CMC	OK	0	No Change	
B3	20	BM400B + CMC	OK	0	No Change	
B4	20	BM400B + CMC	OK	1	No Change	
35	6	BM400B + CMC	OK	0	No Change	
B6	40	BM400B + CMC	OK	0	No Change	
B7	20	BM400B + CMC	OK	0	No Change	
BB	20	BM400B + CMC	NG		or Carange	
B9	20	BM400B + CMC	OK	8	Changed	
B10	20	BM400B + CMC	OK	Š	No Change	
B11	20	BM400B + CMC	OK	0	No Change	

## (Evaluation)

[0186] Evaluations were carried out for the produced porous film and the lithium ion secondary battery completed in the above, as in the following. The results are shown in Tables 3 to 4.

## [Adhesiveness of Porous Film]

[0187] After applying on the positive electrode, the negative electrode, or the separator and then drying, the porous film thus obtained was observed visually immediately after the formation. For those showed a trace of chips, cracks, or separation are marked as "NG" in Table 3.

[Appearance of Negative Electrode]
[0188] After applying the raw material paste for porous

[0188] After applying the raw material paste for porous film on the negative electrode, the paste was dried and the condition of the negative electrode immediately after the formation of the porous film was observed visually. For those showed defects such as size changes are marked as "changed" and others are marked as "changed" in Table 3.

## [Flexibility of Porous Film]

[0189] Upon winding the positive electrode and the negative electrode interposing the sheet separator, the condition of the porsus film formed on any one of the positive electrode, the negative electrode, and the superator was observed visually, mainly the condition near the winding core. For each battery, [10] pieces of the wound electrode plate groups were made, and the numbers of the electrode plate group in which chips, cracks, or separation occurred on the winding are shown in Table 3.

### [Battery Designed Capacity]

[0190] For a diameter of 18 mm of the battery case, the diameter of the wound electrode plate groups were set as 16.5 mm focusing on easiness of insertion. In this case, the battery designed capacity was obtained from positive electrode weight, by setting the capacity per 1 g of the positive electrode active material as 142 mAh. and shown in Table 4.

## [Charge and Discharge Characteristics of Battery]

[0191] For the completed batteries comprising the electrode plate groups without chips, cracks, or separation due to the winding, preliminary charge and discharge were carried out twice, and the batteries were stored for 7 days under an atmosphere of 45° C. Afterwards, under an atmosphere of 25° C., the following two patterns of charge and discharge were carried out.

(1) The First Pattern

[0192] Constant Current Charge: 1400 mA (End Voltage 4.2 V)

[0193] Constant Voltage Charge: 4.2 V (End Current 100 mA)

[0194] Constant Current Discharge: 400 mA (End Voltage 3  $\mathrm{V}$ )

(2) The Second Pattern

[0195] . Constant Current Charge: 1400 mA (End Voltage 4.2 V)

[0196] Constant Voltage Charge: 4.2 V (End Current  $100~\mathrm{mA})$ 

[0197] Constant Current Discharge: 4000 mA (End-voltage 3 V)

 $[\,0198\,]$  . The charge and discharge capacities at this time are shown in Table 4.

[Nail Penetration Safety]

[0199] For those batteries completed the evaluation of charge and discharge characteristics, the following charges were carried out at an atmosphere of 20° C.

[0200] Constant Current Charge: 1400 mA (End Voltage 4.25 V)

[0201] Constant Voltage Charge: 4.25 V (End Current 100 mA)

[0202] The batteries after the charging were penetrated with an iron-made round nail with a diameter of 2.7 mm under an atmosphere of 20°C, with the speed of 5 mm/second or 180 mm/second from the side face of the battery, and best generation at the time was observed. The temperatures reached after 1 second and after 90 seconds at the penetration point of the battery are shown in Table 4.

TARLE 4

					Battery			
						Nail Penetr	ation Safety	
		Charge and Discharge Characteristics			Nail Speed 5 mm/s		Nail Speed 180 mm/s	
	Designed		400 mAh	4000 mAh	Temperate	ire Reached	Temperati	ire Reached
Battery No.	Capacity (mAh)	Charge (mAh)	Discharge (mAh)	Discharge (mAh)	After I sec.	After 90 sec.	After 1 sec.	After 90 se
Al	1944	1941	1938	1819	72	91	70	88
A2	2008	2010	2006	1879	76	89	69	93
A3	2001	1999	1996	1881	71	92	74	94
A4	1943	1941	1939	1821	74	94	72	89
A5	1872	1875	1874	1759	68	88	76	90
A6	1801	1799	1796	1688	69	96	69	96
A7	1729	1730	1728	1622	77	90	70	91
A8	2126	2124	2119	1989	72	91	73	94
A9	2094	2090	2086	1962	70	97	74	87
A10	2014	2016	2012	1902	65	93	72	95
AH	1873	1874	1870	1760	74	96	71	90
A12	1800	1797	1792	1679	71	89	73	94
A13	1942	1940	1937	1784	66	88	69	91
A14	1944	1939	1938	1805	70	96	72	89
A15	1939	1938	1935	1819	70	94	68	89
A16	1944	1942	1939	1823	73	91	73	88
A17	1941	1940	1937	1819	69	88	74	92
A18	1943	1942	1940	1811	72	90	71	93
A19	1946	1944	1941	1799	70	93	69	90
B1	2015	2014	2003	1888	146	-	138	
B2	1944	1939	1935	1812	81	151	69	93
B3	2010	2011	2008	1887	78	139	77	136
B4	1584	1581	1574	1386	73	94	7.5	96
B5	2157	2151	2146	2010	79	144	81	149
B6	1656	1653	1649	1488	20	87	71	90
B7	1940	1848	1822	1472	71	95	73	94
B8								
By	1938	1935	1929	1806	88	146	76	100
B10	1942	1940	1936	1810	78	141	73	94
B11	1938	1937	1934	1816	146	-	142	

#### (Results Observation)

[0203] In the battery B1 having no porous film, the heat generation after I second was notable regardless of the nail penctration speed. On the other hand, in the batteries A1 and A2, in which the porous film was formed on the positive electrode or the negative electrode, the heat generation after the nail penctration was suppressed greatly.

[0204] As all the batteries after nail penetration tests were disassembled and checked, it was found that in all batteries, the sheet separator was melotd in a widespread area. However, in the batteries AI and A2, it was found that the porous film kept its original shape, Based on this, it can be thought that when the procus film has enough host resistance, the abort crevit coverred after the anil penetration will not destroy the lith structure, will suppose the expansion of the other crevit period. The anil will prove the excessive generation of short crevit part, and will prove the excessive generation of

[9295] On the other hand, in the battery B2 in which the provises film was formed on the sheet separator, it was found that the heat generation is accelerated when the nail penetration speed is slow. As the batteries of the battery B2 were disassemibled, it was found that provise film also deformed with the above mentioned meding of the separator. This is probably because when the base substrate which supports when the separator and when the separator fundamental than the separator of the separ

[0206] Herein, characteristics of the nail penetration test, which is a substitutional evaluation for the internal rocticuit, and the data interpretation were explained in detail. First, regarding causes of the heat generation by the present penetration test, following can be explained from the past we obtained results. When a positive electrode and a special content of the content of a both circuit by a nail penetration, the Joule heat is generated. Then, a material (special) with a low heat resistance is melted by the Joule heat to form a strong short circuit portion. As a result, the Joule heat is continued to be generated, and the temperature of the positive electrode is increased to the temperature range (160° C. or more) where the positive electrode becomes thermally unsable. But hem alternavay is thus caused.

[9207]. Also, when the nail penetration speed is decreased, an acceleration of a local heat generation was observed. This is probably because when the short circuit area is limited by slowing the nail penetration speed per a unit time, a considerable amount of heat concentrates on the limited portion, to accelerate the temperature to reach the temperature range where the positive electrode becomes thermally unstable.

[9208] On the other hand, when the mil penetration speed is increased to expand the short circuit area per a unit time, the heat is dispersed to a greater area, thereby making it difficult for the positive electrode to reach the temperature range where it becomes thermally unsubstance.

[0209] Currently, in various uses, the safety standard for the lithium ion secondary battery is becoming strict. Under such circumstances, the present invention which can suppress the thermal runaway is highly practical, regardless of the nail penetration speed (short circuit state).

[0210] As to the thickness of the porous film, in the battery B4 having an excessively thick porous film, designed capacity decreased and capacity under the high-rate discharge decreased, due to shortened length of the electrode plate forming the electrode plate group. Therefore, in order to embody the effects of the present invention sufficiently, it is desirable that the thickness of the porous film is 0.5 to 20 µm.

[9211] In the battery B6 which had the abest separation with an excessive thickness, the designed capacity declined to a greater degree, and the capacity under the high-rate dischange declined, due to the shonened length of the electrode plate which formed the electrode plate group. Therefore, in order to embody the effects of the present invention fully, it is desirable that the thickness of the separator is 30 µm or below. However, in order to suppress the heat generation involved with the melting of the separation of the control of the thickness is \$m\$ or more.

[0212] The filler content in the porous film is explained

[0213] In the battery B7 in which the filler content relative to the filler and the first binder in total was small (binder amount was lange), a capacity decline under the high-rate discharge could be seen. This is probably because the ion conductivity of the promus film declined due to the excessive first binder and insufficient gaps among the filler particles.

[0214] In the battery B9 which used CMC and in the battery B10 which used PVDF, as the first binder in the porous film, the beat generation was not suppressed when the nail penetration speed was slowed. When these batteries were disassembled and checked, it was found that the porous film deformed in addition to the separator.

[9215] In battery B9, CMC (a decomposition temperature of 245° C) was probably burned out due to the Joule heat of the short circuit, causing the damages to the athesiveness of the porous film. Also, in the battery B10, the deformation of the porous was caused probably due to the melling of PVDT (a crystallina melting point of 174° C). Additionally, in both cases, a strong short circuit portion was formed by the penetration of the nail and the heat generation probably was not suppressed.

[9216] Therefore, it is necessary to use, for the porous littin, a first binder which is not easily burned out or melect itself, more concretely, at least one kind of binder having a boundary temperature such as exystalline melting point and a decomposition temperature of 250° °C. or more. From the above evaluation results, if, can be understood that a rubber including an explorative least which is amorphous and has a high heat resistance (a heat decomposition temperature of 320° °C.) can be used preferably.

[0217] The nubber including an acylonitrile unit has niber elasticity. This property linucious very advantageously in the wound-type electrode plate group formation. For example, in the battery A4 in which the binder had the rubber elasticity, the porous film after the winding kept into form sufficiently, and there was no defect. On the other, in the battery B9 and B10, the flexibility evaluation of the prorusa film above poor results.

[0218] Also, in the battery B9, the defects in appearance due to the deformation of the negative electrode were shown after the formation of the porous film. As is mentioned above this is probably the results of the asvelling of the thickners in the negative electrode by water included in the porous film before drying, In order to avoid such tow yields, as water-insoluble first indired should be used for the protus film, and water should not be used as a dispersion medium of the raw material paste for protus film. More generally, a dispersion medium different from the dispersion medium used in the raw menerial paste for protus printing electrode material mixture layer (negative electrode paste) should be used to form the protus film.

[0219] In the battery Al Kin which titunia was used instead of abunitum as the filter, it was confirmed that titunia neahieved various functions almost same as aluminas. On the other band, in the battery Bl II which used polyethylene beads (Pt beads), the level of the nail penetration safety was almost the same level with the battery Bl II which used the persons film irrom the above, it can be seen that the persons film enamed fully having just the same level with the rationary and the filter having just the same level of blue rationary was used. Therefore, it is desirable that for the filler integration voide is selected.

[0220] The structure of the negative electrode is explained in the following.

[0221] As shown by the battery DT, when PVDF was used in the negative electrode the secund hinder content in the negative electrode was ineviably increased, thereby decreasing lithium ion acceptability of the negative electrode to cause a gradual decrease in the charge capacity. Additionally, beads on the property of PVDF, the negative electrode plate was hardened, becoming unable to make the most of the flexibility of the proposition. Herefore, it is desirable that a binder, like SIR which has rubber elusticity and which can give sufficient addiscipancies to the negative electrode material nixture layer by a small amount is used in combination with a water-soluble binder (CMC or the

## EXAMPLE 3

#### Battery C.

(a) Preparation of Raw Material Paste of Porous Film

[9222] A raw material passe for purous film was prepaude by mixing 970 g of AKP90 (cashumian with a median diameter of 0.1 to 0.3 mm) as a filler manufactured by Sumition Chemical (C. 1.4d, 375 g of BM-7201 manufactured by ZEON Corporation (NMP solution containing & wt/s of the first rubber including a polyper/pointiel chain as a first binder), and an appropriate amount of NMP in the double-arm kneeds.

[0223] Properties of BM-720H are shown in the following.

[0224] <1> Crystalline Melting Point: None (amorphous)

[0225] <2> Decomposition Temperature: 320° C.

[0226] <3> Affinity with Water: Water-Insoluble

[0227] The properties of AKP50 are shown in the following.

[0228] <1> Surface Alkalinity: pH=9

[0229] <2> BET Specific Surface Area: about 10 m<sup>2</sup>/g

[0230] <3> Heat Resistance: 250° C. or more

[0231] As shown in the above, since alkalinity of AKP50 was pH=9, it was confirmed that AKP50 had an alkaline site on its surface.

[0232] Also, it is known that AKP50 is  $\alpha$ -atumina, and its heat resistance is known as 250° C, or more.

(b) Eabrication of Positive Electrode

[0233] A positive electrode material mixture paste was prepared by mixting 3 kg of lithium coshaltnet, 1 kg of PVDF/31230 (WhiP Solution constituting 12 wt % of PVDF) manufactured by Kureha Chemical Industry Co. 1 fd. as a binder. 99 g of a seek/ten black, and an appropriate amount of NMIs with a double-orn kneader. This paste was applied and of the properties of the pr

#### (c) Preparation of Negative Electrode

[9234] A negative electrode material mixture poste was propered by mixing 3 kg of artificial graphite, 75 g of BM-400B (squeous dispersion including 40 w; 46 of sydebustiene copolymer as a second binder (second rubber particle)) manufactured by ZEON Corporation. 30 g of CMC as a water-soubhe polymer, and an appropriate amount of water with a double-orm kneeder. This paste was applied on a copper foil with a thickness of 10 µm, and rotted after drying, to form a negotive electrode material mixture layer. At this time, the electrode plate comprising the copper foil and the material mixture layers was set as 180 µm. Afterland the material mixture layers was could be inserted into the can better one of the cylorical battery (product number), to obtain a negative electroda buton.

## (d) Formation of Porous Film

[0235] The raw material paste for porous film was applied on both sides of the positive electrode hoop, and then dried, to form a porous film having a thickness of 5 µm per side adhered to the positive electrode surface.

## (e) Preparation of Electrolyte

[0236] The electrolyte was prepared by dissolving LiPF<sub>6</sub> in a mixed solvent of ethylene carbonate (EC), dimethyl carbonate (DMC), and methyl ethyl carbonate (MEC) with a volume ratio of 2:3:3, to give a concentration of 1 mol/L, and further adding 3 wt % of vinylene carbonate (VC)

## (f) Fabrication of Battery

[9237] The above positive electrode and the negative electrode were cut to give a predetermined length, respectively, and were wound interposing a sheet separator with a thickness of 20 µm comprising a microporous film made of the composition of the composition of the composition of the case the composition of the case was sealed. A cylindrical-shape lithium ion secondary battery CI was this made.

Battery D1

[0238] A battery D1 was made in the same manner as the battery C1, except that the porous film was not formed on both sides of the positive electrode hoop.

Battery D2

[0239] The mw material paste for porous film was applied on both sides of the separator comprising polythylasemade microporous film, and then dried, to form a poreus film having a thickness of 5 µm per side adhered to the separator surface. Battery 10 µm sen adia in the same manner as the battery 11), except that the separator having the porous film thus obtained was used:

Battery C2

[0240] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 0.5 µm per side adhered to the negative electrode surface. Battery C2 was made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was used.

Battery C3

[0241] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 1 µm per side adhered to the negative electrode surface. Battery C3 was made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was used.

Battery C4

[0242] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 5 µm per side adhered to the negative electrode surface. Battery C4 was made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was

Battery C5

[0243] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 10 µm per side adhered to the negative electrode surface. Battery CS was made in the same manner as the battery DI, except that the negative electrode having the porous film thus obtained was used.

Battery C6

[0244] The raw material paste for provos film was applied on both sides of the negative electrode hoop, and then dried, to form a prorus film having a thickness of 15 µm per side adhered to the negative electrode surface. Battery C6 was made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was used.

Battery C7

[0245] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film having a thickness of 20 µm per side adhered to the negative electrode surface. Battery C7 was

made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was used.

Battery C8

[0246] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a prouse film having a thickness of 30 µm per side adhered to the negative electrode surface. Battery CS was made in the same manner as the battery D1, except that the negative electrode having the porous film thus obtained was

Battery C9

[0247] A battery C9 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 8 µm.

Battery C10

[0248] A battery C10 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 10 µm.

Battery C11

[0249] A battery C11 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 15 µm

Battery C12

[0250] A battery C12 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 25 µm.

Battery C13

[0251] A battery C13 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 30 µm.

Battery C14

[0252] A battery C14 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the thickness of the sheet separator was set as 40 µm.

Battery C15

[9253] A battery C15 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 30 wt %.

#### Battery C16

[0254] A battery C16 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 50 wt %.

## Battery C17

[0255] A bottery C17 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 70 µm (%).

#### Battery C18

[0256] A battery C18 in which a thickness of the porous film adhered to the negative electrode surface was 5  $\mu$ m per side thereof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 90 at 9%.

#### Battery C19

[0257] A battery C19 in which a thickness of the porous libra adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 95 w 4%.

#### Battery C2

[9288] A battery C20 in which a thickness of the porous film adhered to the negative electrode surface was 5 jumper side theroof was made in the same manner as the battery C4, except that the filler content relative to the total of the filler and the first binder was set as 99 wt %.

#### Battery D3

[0259] A buttery D3 in which a thickness of the porous lim adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery D4. except that a water-soluble CMC was used instead of BM-72011 as the first binder to prepare the raw material paste for perous film. Since CMC was used as the first binder, in Battery D3, as a dispersion medium of the raw material paste, water was used instead of DMP. CMC does not have a crystalline melting point, is amorphous, and has a decomposition temperature of 245° C.

## Battery D4

[9260] A battery D4 in which a thickness of the porous lift adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery D4. except that water-insoluble PDF was used as the first binder instead of BM-7201 to prepare the raw material paste for porous film. A ceystalline melting point and a decomposition temperature of PVDF are 174° C. and 360° C. respectively.

#### Battery C21

[0261] A battery C21 in which a thickness of the porous film adhered to the negative electrode surface was 5  $\mu$ m per side thereof was made in the same manner as the battery C4, except that, instead of AKP50 ( $\alpha$ -alumina having a median diameter of 0.1 to 0.3  $\mu$ m), titania (titanium oxide) having the same median diameter was used

[0262] For the titania, TA300 (anatase-type) manufactured by FUJI TITANIUM INDUSTRY CO., LTD. was used. The BET specific surface area of the titania was 8 m²/g, and the titania surface had an alkalinity of pH-8. The heat resistance of polyethylene beads was below 250° C., the BET specific surface area was 10 m²/g, and the pH of beads surface ware

#### Battery C22

[928] A battery (22 in which a thickness of the porsus him addrered to the negative electrode surface was 5 tim per side thereof was made in the same maner as the state of the except that a linear type PPS (psh place) ten salidation; Cafan diameter of 0.5 mm) manufactured by DANIPPON INK AND CHEMICA (3. N.CORPORATED was used to the of AKPS). The PIS surface had an alkalinity of pl. 1e.7 at 6. the BIT specific surface area was 7.4 m<sup>2</sup>/g, and the PIS had an excellent heat-resistance over 250° C. (melting point of 280° C.).

#### Battery C23

[9264] A battery C23 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same namer as the battery C4, except that AA2 (alumina Z) manufactured by Sumitomo Chemical Co., Ltd. was used instead of ARP50. The AA2 surface had an alkalinity of p11=9, and the BET specific surface area of 10 m<sup>2</sup>/<sub>2</sub>/<sub>2</sub>.

#### Battery C24

[0265] A battery C24 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that the titania (titania Y) with the surface alkalinity of pH-8, and the BET specific surface area of 1.0 m<sup>2</sup>/g was used instead of AKP50.

#### Battery C25

[0266] A battery C25 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm estimated in the same manner as the battery C4, except that AA2 (albumina Y) manufactured by Sumitoma Chemical Co. 1.d. was used instead of AKP50. The AA2 surface had an alkalinity of pH=9, and the BET specific surface area of 0.9 m<sup>2</sup>/a.

## Battery C26

[0267] A battery C26 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4, except that titania (titania X) with the surface alkalinity of p11–8, and the BET specific surface area of 0.9 m<sup>2</sup>/g was used instead of AKPS0.

## Battery D5

[9268] A battery DS in which a thickness of the porous film adhered to the negative electrical surface was 5 μm per side thereof was made in the same manner as the battery (\*4, except that crosslinked-type PS (nopyhenyber) esufficie, median diameter of 0.5 μm) was used instead of A&P50. The BETT specific surface area of the crosslinked-type PSS was 7.4 m/g, same as the above, and the pH of the surface theored was 6. Battery D6

[0269] A battery D6 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery C4.

[0271] The configuration of the porous film is summarized and shown in Table 5. Additionally, the thickness of the sheet separator and the kinds of the second binder used for the negative electrode are summarized and shown in Table

TABLE 5

				Porous Film										
						First Buider								
		Frlm	Fdl	er		Crystalline Melting	Decomposition							
Battery No.	Adhered Portion	Thickness (µm)	Kind	Content (wt %)	Kind	Point (° C.)	Temp.							
CI	Pos. F.lec	5	AKP50	97	BM720H	None	320							
C2	Neg. Elec.	0.5	AKP50	97	BM720H	None	320							
23	Neg. Elec.	Ţ	AKP50	97	BM720H	None	320							
C4	Neg. Elec.	5	AKP50	97	BM720H	None	320							
C5	Neg. Elec.	10	AKP50	97	BM720H	None	320							
C6	Neg. Elec.	1.5	AKP50	97	BM720H	None	320							
27	Neg. Elec.	20	AKP50	97	BM720H	None	3.20							
.8	Neg Hec.	30	AKP50	97	BM720H	None	320							
.9	Neg. Elec.	5	AKP50	97	BM720H	None	320							
210	Neg. Elec.	5	AKP50	97	BM720H	None	320							
11	Neg. Elec.	5	AKP50	97	BM720H	None	320							
12	Neg. Elec.	5	AKP50	97	BM720H	None	320							
213	Neg. Elec.	5	AKP50	97	BM720H	None	320							
14	Neg. Hec.	5	AKP50	97	BM720H	None	320							
15	Neg. Elec.	5	AKP50	50	BM720H	None	3.20							
.16	Neg Elec.	5	AKP50	70	BM720H	None	320							
217	Neg. Elec.	5	AKP50	90	BM720H	None	320							
18	Neg Elec.	5	AKP50	95	BM720H	None	320							
19	Neg. Elec.	5	AKP50	99	BM720H	None	320							
220	Neg. Elec.	5	AKP50	30	BM720II	None	320							
221	Neg. Elec.	5	TA300	97	BM720H	None	320							
22	Neg. Elec.	5	Alkaline PPS	97	BM720H	None	320							
23	Neg. Elec.	5	Alumina Z	97	BM720H	None	320							
24	Ncg. Elec.	5	Titania Y	97	BM720H	None	320							
:25	Neg. Flec.	5	Alumina Y	97	BM720H	None	320							
26	Neg Elec.	5	Titania X	97	BM720H	None	320							
01	None													
02	Separator	5	AKP50	97	BM720H	None	320							
03	Neg. Elec.	5	AKPS0	97	CMC	None	245							
)4	Neg. Elec.	5	AKP50	97	PVDF	174	360							
)5	Neg Flee.	5	Acidsc PPS	97	BM720H	None	320							
36	Neg. Islec.	5	Silica	97	BM720H	None	320							
<b>D</b> 7	Neg. Elec.	5	Alumina X	97	BM720H	None	320							

except that a silica powder was used instead of AKP50. For the silica powder, Sunsphere NP30 manufactured by ASAHI (HASS CO, LTD, was used. The BET specific surface area of the silica powder was 40 m<sup>2</sup>/g, and the pH of the surface of the silica powder was 6.

#### Battery D7

[9279] An alumina (alumina X) with the BET specified surface area of 10 m<sup>2</sup>/g, and the surface of pHef-98 section instead of AKPS0. Additionally, in fibricating the negative electrode, instead of BM400B as a binder B and CMC as a tluckence, 8 wt% of PVDF relative to the artificial graphia was used. Herein, for the PVDF. PVDF#1320 used to the binder of the positive electrode was used. Other than the above, a battery Dri a which the porus film adhered the negative electrode surface has a thickness of 5 im per side thereof was made in the same manner as the battery C4.

## [0272]

TABLE 6

			Porc	nas Film	
Bat- tery No	Separator Thickness (jim)	Negative Electrode Binder	Ad- hesive- ness	Flexibility (No. of Defects)	Negative Electrode Appearance
C1	20	BM400B + CMC	OK	0	No Change
C2	20	BM400B + CMC	OK	0	No Change
C3	20	BM400B + CMC	OK	0	No Change
C4	20	BM400B + CMC	OK	0	No Change
C5	20	BM400B + CMC	OK	0	No Change
C6	20	BM400B + CMC	OK	0	No Change
C7	20	BM400B + CMC	OK	0	No Change
C8	20	BM400B + CMC	OK	1	No Change
C9	8	BM400B + CMC	OK	0	No Change
CIU	10	BM400B + CMC	OK	0	No Change

Separator

(jan)

25 30

20

20

20

20

BM400B + CMC OK

BM400B + CMC OK

BM400B + CMC OK

BM400B + CMC OK

BM400B + CMC

BM400B + CMC ОК

BM400B + CMC OK

BM400B + CMC OK

BM400B + CMC OK

tery No Thickness

C13 C14

C18

C19 20

C20 20

C22 201

C24

C25 20

C26 20

D1 20 BM400B + CMC BM400B + CMC

D2 20 No Change

Porous Film Flexibility Negative Ad. Negative Electrode hegive. (No. of Electrode Binder ness Defects) Appearance BM400B + CMC No Change OK BM400B + CMC OK No Change BM400B + CMC OK No Change BM400B + CMC BM400B + CMC OK No Change OK No Change BM400B + CMC OK No Change BM400B + CMC OK No Change

OK

OK

TABLE 6-continued

IABLE 6-continued

Bat- tery No			Porc		
	Separator Thickness (µm)	Negative Electrode Binder	Ad- hearve- ness	Flexibility (No. of Defects)	Negative Flectrode Appearance
D3	20	BM400B + CMC	OK	8	Changed
D4	20	BM400B + CMC	OK	5	No Chanse
D5	20	BM400B + CMC	OK	3	No Change
D6	20	BM400B + CMC	OK	4	No Change
D7	20	PVDF	OK	2	No Change

#### (Evaluation)

[0273] For the above produced porous film and the completed lithium ion secondary battery, porous film adhesiveness, negative electrode appearance, porous film flexibility, battery designed capacity, charge and discharge characteristics of the battery, and the nail penetration safety were evaluated in the same manner as Example 2. The results are shown in Tables 6 to 7.

TABLE 7

					Battery					
					Nail Penetration Safety					
		Charge and Discharge Characteristics				Speed nm/s	Nail Speed 180 mm/s			
	Designed	Designed	400 mAh 4000 mAh		Temperati	ire Reached	Temperatu	re Reached		
Battery No.	Capacity (mAh)	Charge (mAh)	Discharge (mAh)	Discharge (mAh)	After 1 sec.	After 90 sec. (° C.)	After 1 sec.	After 90 se		
CI	1944	1941	1938	1819	72	91	70	88		
C2	2008	2010	2006	1879	76	89	69	93		
C3	2001	1999	1996	1881	71	92	74	94		
C4	1943	1941	1939	1821	74	94	72	89		
C5	1872	1875	1874	1759	68	88	76	90		
C6	1801	1799	1796	1688	69	96	69	96		
C7	1729	1730	1728	1622	77	90	70	91		
C8	1584	1581	1574	1386	73	94	75	96		
C9	2126	2124	2119	1989	72	91	73	94		
C10	2094	2090	2086	1962	70	97	74	87		
C11	2014	2016	2012	1902	65	93	72	95		
C12	1873	1874	1870	1760	74	96	71	90		
C13	1800	1797	1792	1679	71	89	73	94		
C14	1656	1653	1649	1488	70	87	71	90		
C15	2016	1988	1961	1737	66	88	69	91		
C16	2015	1999	1989	1811	70	96	72	89		
C17	2017	2019	2015	1893	70	94	68	89		
C18	2014	2015	2009	1883	73	91	73	88		
C19	2015	2015	2010	1886	69	88	74	92		
20	2017	1848	1822	1472	71	95	73	94		
C21	2014	2012	2005	1880	72	90	71	93		
:22	2014	2010	2002	1874	71	89	69	88		
C23	1944	1940	1938	1822	25	93	73	91		
24	1944	1941	1937	1819	75	95	74	91		
C25	1942	1923	1889	1769	78	92	7.5	95		
26	1939	1919	1881	1762	77	98	74	90		
D1	2015	2014	2003	1888	146		138			
D2	1944	1939	1935	1812	81	151	69	93		
D3	2012	2014	2004	1880	88	146	76	160		
D4	2015	2016	2011	1896	78	141	73	94		
DS	2013	1837	1811	1789	83	149	82	147		
D6	2014	1796	1746	1704	87	154	88	151		
D7	2016	1961	1958	1819	70	93	69	90		

#### (Results Consideration)

[0274] First, in the battery D1 having no prouts film regardless of the null potentiation speed, the heat generation after 1 second was notable. As opposed to this, in the batteries C1 and C2 in which the prorous film was formed on the positive electrode or the negative electrode, the heat generation after the null potentiation was suppressed greatly. When all the batteries were disassembled and checked after the all penetration test, in all batteries, the sheet superator the sail penetration test, in all batteries, the sheet superator the batteries C1 and C2, the prowns lift kept its original shape.

[0275] On the other hand, in the battery D2 in which the porous film was formed on the sheet separator, it could be seen that the heat generation was accelerated when the nail penetration speed was slow. When the battery D2 was disassembled and checked, it was confirmed that the porous film deformed with the melting of the separator.

[0276] With regard to the thickness of the porous film, in the battery C8 with an excessive thickness of the porous film, due to the shortened length of the electrode plate which formed the electrode plate group, designed capacity was decreased, and the capacity under the high-rate discharge was declined.

[0277] In the battery C14 with an excessive thickness of the sheet separator, due to the shortened length of the electrode plate which formed the electrode plate group, designed capacity was greatly decreased, and the capacity under the high-rate discharge was declined.

[9278] Then, in the battery C20 in which the filler content relative to the filler and the first binder in total was small (binder amount was large), the capacity decline under the high-rate discharge could be seen. This is probably because the ion conductivity of the porous film declined due to the excessive first binder and insufficient gaps among the filler particles.

[0279] In the battery D3 in which CMC was used and in the battery D4 in which PVDF was used, as the first binder in the portous film, the heat generation could not be suppressed when the nail penetration speed was decreased. When these batteries were disassembled and checked, it could be confirmed that not only the separator, but the portous film deformed as well.

[0280] Also, although in the battery C4 in which the binder had rubber elsaticity, it was shown that the porous film after winding kept its shape sufficiently and had no defects, an the batteries D3 and D4, the results of the evolution for the porous film flexibility were poor. Further, in the battery D3, the appearance defects due to the negative electrode deformation were seen after the formation of the porous film. This is probably the result of the swelling of the thickner in the negative electrode by water included in the porous film. This ordying, as mentioned above.

[0281] Also, in the battery C21 in which titania was used instead of alumina as the filler, it was confirmed that titania achieved various functions almost same as alumina.

[0282] Next, as shown by the battery D7, when PVDF was used for the second binder of the negative electrode, the amount of the second binder inevitably increased, to decrease lithium ion acceptability of the negative electrode

and to gradually decrease the charge capacity. Also, based on the properties of PVDF, the negative electrode plate hardened

[0283] Also, with regard to the speed of the electrolyte injection at the time of battery labrication, the battery C25 unjection at the time of battery labrication, the battery C25 unique labrication to the battery C25 unique labrication time. Also the battery C24 taking 16 times more injection time. On the other hand, the tipication speeds in the battery C34 unique labrication time. On the other hand, the tipication speeds in the battery C23 were not much different and the injection speeds in the battery C23 and in the battery C24 and in the battery C23 and in the battery C24 were not much different as well. From the above, it can be seen that the ease of the electrolyte injection changes greatly depending on the BET specific surface area of the electrolyte injection changes greatly depending on the BET specific surface area of the eight with Og m<sup>2</sup>g as the borderline. This, it can be seen that the BET specific surface area of the great of the other with Og of the other with Og of the Og o

## EXAMPLE 4

## Battery B1

## (a) Preparation of Raw Material Paste of Porous Film

[9284] A raw material pasts for porous film was propared by mixing 970 g of an anists-type tinnium oxide powder (median diameter 0.09 µm) which was classified by a sive to give the maximum particle size of 0.1 µm, 375 etc. BM-720H (NMP solution containing 8 wt % of a maherincluding an arryloyaltite uit, as a first bidnerly manieratured by Zeon Corporation, and an appropriate amount of NMP in a double-sum kneader.

[0285] The properties of the titanium oxide powder used in this Example are shown below.

[0286] <1> Surface Alkalinity: pH=8

[0287] <2> Heat Resistance: 800° C. or more

[0288] <3> Particle Size: The Maximum Particle Size 0.1 um

#### (b) Fabrication of Positive Electrode

[9289] A positive electrode material mixture paste was perpeared by mixing 3 kg off lithium cobaltate, 1 kg of PVDF#1330 (N-methyl-2-pyrrolidone (NMP) solution including 12 wt w6 of PVDF# namificatured by Kurcha Chemical Industry Co., Ltd. as a binder, 90 g of acceybre back as a conductive agent, and an appropriate amount of NMP, with a double-arm kneader. This paste was applied on a ulaminum foil with a thickness of 15 µm, and rolled after drying to form a positive electrode material mixture layer. At the state, the thickness of the electrode plate comprising the comparison of the properties of the

## (c) Fabrication of Negative Electrode

[0290] A negative electrode material mixture paste was prepared by mixing 3 kg of artificial graphite, 75 g of 8M-400B (aqueous dispersion including 40 wt % of stylenebutadiene copolymer (rubber particles) as a second binder) manufactured by ZEON Corporation, 30 g of CMC as a thickener, and an appropriate amount of water with a doubleh arm kneader. This paste was upplied on a copper foil with thickeness of 10 mm, and rolled after drying, to form a negative electrod meteral mixture layer. At this time the thickness of the electrode plate comprising the copper foil and the material mixture layers was et as 180 µm and and the material mixture layers was set as 180 µm and wards, the electrode plate was slit to give a width whiteir could be inserted into the can battery case of the cylindrical battery (product number 18650), to obtain a negative electrode hoop.

#### (d) Formation of Porous Film

[0291] The raw material paste for porous film was applied on hoth sides of the positive electrode hoop, and then dried, to form a porous film having a thickness of 5 µm per side adhered to the positive electrode surface.

## (e) Preparation of Electrolyte

[0292] An electrolyte was prepared by dissolving LiPF<sub>o</sub> in a mixed solvent of ethylene carbonate (EC), dimethyl carbonate (DMC), and methyl ethyl carbonate (MEC) with a volume ratio of 2:3:3, to give a concentration of I moVL and further adding 3 wt % of winglene carbonate (VC) thereto.

#### (f) Fabrication of Battery

(9293) The above positive electrode and the negative electrode were tot give a predetermined length, respectively, and were wound interposing a sixeet separator with a thickness of 20 jun comprising a microporous film made of polyothylene, and then were inserted into the battery case. Then, 5.5 g of the above electrolyte was weighed and injected into the battery case, and the opening of the case was sealed. A cylindrical-shape 18650 lithium ion secondary battery El was thus made.

#### Battery F1

[0294] A battery F1 was made in the same manner as the battery E1, except that the porous film was not formed on both sides of the positive electrode hoop.

#### Battery F2

[0295] The raw material paste for porous film was applied on both sides of the sheet separator, and then dried, to form a porous film with a thickness of 5 µm per side adhered to the separator surface. A battery F2 was made in the same manner as the battery F1, except that the separator having the porous film thus obtained was used.

#### Battery E2

[0296] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 0.5 µm per side adhered to the negative electrode. A battery E2 was made in the same manner as the battery F1, except that the negative electrode having the porous film thus obtained was used.

#### Battery 1/3

(9297) The raw material paste for porous lifta was applied to both sides of the negative electrode hoop, and then dried, to form a porous lifta with a thickness of 1 µm per side adhered to the negative electrode surface. A battery E3 was made in the same manner as the battery F1, except that the negative electrode having the porous film thus obtained was used.

## Battery E4

[0298] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 5 µm per side adhered to the negative electrode surface. A battery F3 was made in the sume manner as the battery F1, except that the negative electrode having the porous film thus obtained was

## Battery E5

[0299] The raw material paste for porous film was applied no both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 10 µm per side adhered to the negative electrode surface. A hattery F5 was made in the same manner as the battery F1, except that the negative electrode having the porous film thus obtained was used.

#### Battery E6

[0300] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 15 µm per side adhered to the negative electrode surface. A battery F6 was made in the same manner as the battery F1, except that the negative electrode having the porous film thus obtained was used.

## Battery E7

[0301] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 20 µm per side adhered to the negative electrode surface. A battery E7 was made in the same manner as the battery F1, overpt that the negative electrode having the porous film thus obtained was used.

## Battery E8

[0302] The raw material paste for porous film was applied on both sides of the negative electrode hoop, and then dried, to form a porous film with a thickness of 30 µm per side adhered to the negative electrode surface. A battery E8 was made in the same manner as the battery F1, except that the negative electrode having the porous film thus obtained was used.

#### Battery E9

[0303] A battery E9 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery E4, except that the thickness of the sheet separator was set as 8 µm.

## Battery E10

[0304] A battery E10 in which a thickness of the porous film adhered to the negative electrode surface was 5 μm per side thereof was made in the same manner as the battery F4, except that the thickness of the sheet separator was set as 10 μm.

#### Battery E11

[0305] A battery E11 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery E4, except that the thickness of the sheet separator was set as 15 µm

#### Battery E12

[0306] A battery F12 in which a thickness of the porous film adhered to the negative electrode surface was 5 jum per side thereof was made in the same manner as the battery E4, except that the thickness of the sheet separator was set as 25 ium.

## Battery E13

[0307] A battery E13 in which a thickness of the porous film adhered to the negative electrode surface was 5  $\mu$ m per side thereof was made in the same manner as the battery E4, except that the thickness of the sheet separator was set as 30  $\mu$ m.

#### Battery E14

[0308] A battery IF14 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery IE4, except that the thickness of the sheet separator was set as 40 µm.

#### Battery E15

[0309] A battery E15 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery E4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 50 µm?

## Battery E16

[0310] A battery E16 in which a thickness of the porous film adhered to the negative electrode surface was 5 jun per side thereof was made in the same manner as the battery E4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 50 in %.

## Battery E17

[0311] A battery E17 in which a thickness of the porous film adhered to the negative electrode surface was 5 jun per side thereof was made in the same manner as the battery E4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 70 mt 9.

#### Battery E18

[0312] A battery E18 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery E4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 90 wt %.

#### Battery E19

[0313] A battery £19 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the buttery £4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 95 wt w.

#### Battery E20

[0314] A battery E20 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery E4, except that the content of the titanium oxide powder relative to the total of the titanium oxide powder and the binder was set as 90 up.

#### Battery F3

[0315] A battery F3 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same mammer as the battery F4, except that a watersoluble CMC was used instead of BM-720 as a binder to prepare the raw material paste for porous film. Herein, for the dispersion medium of the raw material paste, water was used instead of NMP.

#### Battery F4

[0316] A bettery F4 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the bartey E4, except that a water-insoluble PVDF (crystalline melting point 174° C.) was used instead of BM-720 as a binder to prepare the raw material passe for porous film.

#### Battery F5

[0317] A battery F 5 in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery 1:4. except that to-diminian with a median diameter of 0.3 µm was used instead of the thanium oxide powder with a median diameter of 0.09 µm. For the α-alumina, AKP50 manufactured by Sumiton Chemical Co., Lid, was used (such alkalinity pH=9, BET specific surface area: about 10 m<sup>2</sup>/g, heat resistance: 250°C c. or more).

#### Battery F6

[0318] In the fabrication of the negative electrode, instead of BM4008 as a second binder and CMC as a thickner. 8 w % of PVDF relative to the artificial graphite was used. and for the dispersion medium of the negative electrode passe. NMP was used. Herein, for the PVDF, PVDF#1320 which was used as a binder of the positive electrode passe. NMP was used. Herein, for the PVDF, PVDF#1320 which was used as a binder of the positive electrode was used. Except for the above, a battery Fe in which a thickness of the porous film adhered to the negative electrode surface was 5 µm per side thereof was made in the same manner as the battery 184.

[0319] The configuration of the porous film is summarized and shown in Table 8. The thickness of the sheet separator and the kind of the second binder used for the negative electrode were summarized and shown in Table 9.

TABLE 8

				Porous	îlm				
					First Binder				
		Film	Fill	ier		Crystalline Melting	Decomposition		
Battery No.	Adhered Portion	Thickness (µm)	Kind	Content (wt %)	Kind	Point (° C.)	Temperature (° C.)		
E1	Positive electrode	5	Titanium oxide	97	BM720H	None	320		
E2	Negative electrode	0.5	Titsnium oxide	97	BM720H	None	320		
E3	Negative electrode	1	Titanium oxide	97	BM720H	None	320		
E4	Negative electrode	5	Titanisun oxide	97	BM720H	None	320		
1,5	Negative electrode	10	litanam oxide	97	BM720H	None	320		
E.0	Negative electrode	15	Titansam oxide	97	BM720H	None	320		
1-7	Negative	20	Titanum oxide	97	BM720H	None	320		
H.R	Negative electrode	30	Totamum oxide	97	BM720H	None	320		
E9	Negative electrode	5	Titanium oxide	97	ВМ720Н	None	320		
E10	Negative electrode	5	Titanrum oxide	97	BM720H	None	320		
EH	Negative electrode	5	Titanium oxide	97	BM720H	None	320		
E12	Negative electrode	5	Titanium oxsde	97	BM720H	None	320		
E13	Negative electrode	5	Titannum oxide	97	BM720H	None	320		
E14	Negative electrode	5	Titannım oxide	97	BM720H	None	320		
E15	Negative electrode	5	Titnniun oxide	30	BM720II	None	320		
E16	Negative electrode	5	Titamum oxide	50	BM720H	None	320		
317	Negative	5	Titanium oxide	70	BM720H	None	320		
:18	Negative electrode	5	Titanium oxide	90	ВМ720Н	None	320		
19	Negative electrode	5	Titanium oxide	95	BM720H	None	320		
:20	Negative electrode	5	Fitanium oxide	99	BM720H	None	320		
31	None		-						
2	Separator	5	Titannum oxide	97	ВМ720Н	None	320		
3	Negative electrode	5	Istanium oxide	97	CMC	None	245		
4	Negative electrode	5	Titanium	97	PVDF	174	360		
5	Negative electrode	5	α- alumina	97	ВМ720Н	None	320		
6	Negative electrode	5	Titanium	97	BM720H	None	320		

[0320]

TABLE 9

		17401							
	Porous Film								
But- tery No.	Separator Thickness (µm)	Negative Electrode Binder	Ad- heave- ness	Flexibility (No of Defects)	Negative Electrode Appearance				
F1	20	BM400B + CMC	OK	0	No Change				
E2	20	BM400B + CMC	OK	0	No Change				
L3	20	BM400B + CMC	OK	0	No Change				
E4	20	BM400B + CMC	OK	0	No Change				
E5	20	BM400B + CMC	OK.	0	No Change				
E6	20	BM400B + CMC	OK.	0	No Change				
E7	20	BM400B + CMC	OK	0	No Change				
E8	20	BM400B + CMC	OK.	1	No Change				
E9	8	BM400B + CMC	OK	0	No Change				
E10	10	BM400B + CMC	OK	0	No Change				
E11	15	BM400B + CMC	OK	0	No Change				
E12	25	BM400B + CMC	OK	0	No Change				
E13	30	BM400B + CMC	OK	0	No Change				
F14	40	BM400B + CMC	OK	0	No Change				
E15	20	BM400B + CMC	OK	0	No Change				
H16	20	BM400B + CMC	OK	0	No Change				
E17	20	BM400B + CMC	OK	0	No Change				
E18	20	BM400B + CMC	OK	0	No Change				
E19	20	BM400B + CMC	OK	0	No Change				
E20	20	BM400B + CMC	OK	1	No Chang				
Fl	20	BM400B + CMC			No Chang				
F2	20	BM400B + CMC	OK	0	No Change				

TABLE 9-continued

			Рого	us Film	
Bal- tery No.	Separator Thickness (jum)	Negative Electrode Binder	Ad- hesive- ness	Flexibility (No. of Defects)	Negative Electrode Appearance
F3	20	BM400B + CMC	ок	8	Changed
F4	20	BM400B + CMC	OK	5	No Change
F5	20	BM400B + CMC	OK	0	No Change
F6	20	PVDF	OK	2	No Change

## (Evaluation)

[0321] For the above produced porous film and the completed lithium ion secondary battery, perous film athesiveness, negative electrode appearance, prours lift in Rixbility, battery designed capacity, charge and discharge characteristics of battery, and the anil penetration safety were evaluated in the same manner as Example 2. The results are shown in Tables 9 to 10.

TABLE 10

					Battery					
					Nail Penetration Safety					
		Charge And Discharge Characteristics			Natl Speed 5 mm/s		Nail Speed 180 nun/s			
	Designed	Designed		400 mAh 4000 mAh		Temperature Reached		re Resched		
Battery No.	Capacity (mAh)	Charge (mAh)	Discharge (mAh)	Discharge (mAh)	After 1 sec. (° C.)	After 90 sec- (° C.)	After 1 sec.	After 90 se (* C.)		
E1	1944	1941	1939	1822	72	90	68	87		
B2	2008	2010	2007	1879	76	89	69	93		
E3	2001	1999	1997	1883	71	91	74	94		
L4	1943	1941	1939	1823	74	92	72	89		
E5	1872	1875	1874	1764	68	88	76 68	96		
E6	1801	1799	1796	1691	69	94	70			
E7	1729	1730	1728	1622	77	89	75	91 96		
E8	1584	1581	1576	1386	73 72	91	73	96		
E9 E10	2126 2094	2124	2119 2086	1989	70	96	72	87		
			2080		65	93	72	94		
EII	2014	2016	2012 1870	1904	74	93	71	99		
E12	1873	1874		1679	71	89	73	90		
E13	1800	1797	1793	1679	71 68	87	71	94		
E14 E15	1656 2017	1653 1848	1649 1822	1486	71	95	73	90		
E15	2017	1988	1961	1737	65	88	69	91		
E16 E17	2016	1988	1981	1812	70	94	72	89		
E17	2015	2019	2015	1812	70	94	68	88		
E18	2017	2019	2015	1884	73	91	73	88		
E20	2014	2015	2009	1888	69	88	74	92		
1-10 1-1	2015	2013	2011	1888	146		138	74		
F2	1944	1939	1935	1812	81	151	69	93		
F3	2012	2014	2004	1881	88	147	76	101		
F4	2012	2616	2011	1897	76	141	73	94		
FS.	1943	1941	1939	1821	74	94	72	89		
F6	1944	1911	1889	1759	70	93	69	90		

#### (Results Consideration)

[9322] In the battery FI having no protous film, the best generation after I second was notable-regardless of the generation speed. As opposed to this, in the batteries EI and IE2, in which the perrous film was formed on the positive electrode or the negative electrode, the heat generation after the nail penetration was greatly suppressed. When all the batteries after the nail penetration test were dissessembled and checked, in all batteries, the sheet separation method is wisconsistent of the state of the penetration of the batteries EI and I2, the pornous film kept its original shape.

[9323] On the other hand, in the battery F2, in which the perous tilm was formed on the separator, it could be sent that the heat generation was accelerated when the nail penetration speed was slow. When the battery F2 was disassembled and checked, it was confirmed that the prorus illin also deformed with the above mentioned melting of the separatior

[0324] With regard to the thickness of the porous film, in the battery ER having a porous film with excessive thickness, since the length of the elsectrode plate forming the electrode plate group was shortened, the designed capacity declined and the capacity at the high-rate discharge declined.

[0325] In the battery E14 having a separator with an excessive thickness, since the electrode plate forming the electrode plate group shortened, the designed capacity greatly declined, and the capacity at the high-rate discharge greatly declined.

[0.326] In the hattery 115, in which the content of tinnium oxide powder relative to the total of the tilinaim oxide powder relative to the total of the tilinaim oxide powder relative to the total of the tilinaim oxide powder and binder was small (binder amount was large), a capacity decline can be seen at the high-rar existence in the second to the second to the second to the second on the second due to the oxec section amount of binder and caused the decline in ion conductivity of the partus film. However, when the content of tilnaimm oxide parture of the second to th

[0327] In the battery F3 in which CMC was used and in the battery F4 in which PVDF was used as the binder in the porous film, the heat generation could not be suppressed when the nail penetration speed was slowed. When these batteries were disassembled and checked, it was confirmed that not only the separator, but the porous film was deformed as well.

[9328] In the hattery F3, defective appearance due to the deformation of the negative electrode could be seen after the formation of the porous film. As is mentioned above, this is probably the results of the swelling of the thickener in the negative electrode due to the water included in the porous film before drying.

[0329] Additionally, in the battery F5 in which calumina was used instead of titanium oxide powder, it was confirmed that titanium oxide achieved various functions almost same as calumina. However, when calumina was used, battery performances varied as the porous film became thinner. Therefore, when the porous film is to be formed thin, titanium oxide is preferably used, instead of calumina. [0330] The reason that battery performances varied as the porous film using c-alumina became thinner was probably that in c-alumina, the gaps having a pore size of 0.1 µm (100 nm) or less cannot be formed uniformly. This implies that a certain degree of thickness is necessary in the case of the porous film using c-alumina.

[0331] As shown by the battery F6, when PVDF was used as the second binder in the negative electrode, safety could be secured. However, the second binder content became inevitably large, and lithium ion acceptability of the negative electrode declined and the charge capacity gradually declined.

#### EXAMPLE 5

Battery G1

(a) Preparation of Raw Material Paste of Porous Film

[9332] A raw material posice for porous film with a viscosity of 48 ps. 20 pm measured by 18-type visconneter (measurement temperature 25° C.) was prepared by mixing 970 g. of alumina with a median diameter of 0.3 µm, 375 g of BM-720H (NMP solution containing 8 wt % of a rubberineluding a polysory-loutifiel chain as a first bindery manfactured by ZEON Corporation, and an appropriate amount of NMP in a double-smi Recader.

#### (b) Fabrication of Positive Electrode

[933] A positive electrode material mixture passe was prepared by mixing 3 kg of linkinu cobaltate. 1 kg of PVDF41320 (NMP solution containing 12 w 1% of PVDF41320 (NMP solution containing 12 NMP, with a coubse-arm kencelor. This peats was applied on an attuminum foil with a thickness of 15 µm, and rolled and redyring, to form a positive electrode material mixture layer. At this time, the thickness of an electrode plate usually a sitting to the property of the prope

## (c) Fabrication of Negative Electrode

[9334] A negative electrode material mixture pasis was prepared by mixed 2 kg of artificial graphitic. 1 kg of vapor growth carbon filher (VGCF), 75 g of BM-400B (sugressor dispersion including 40 wt % of stylene-butudence copolymer (second rubber particle) as a second binder) manufactured by ZION (Corporation, 30 g of CMC as a successful below polymer, and an appropriate amount of water with a couble-arm kaneel. This paste was applied on a coper foil with a thickness of 10 jun; and rolled after drying, to form continued to the control pastern control instruct payr. At this time, the electrode place material mixeus plays At this time, the electrode place was sit to give a width which could be inserted into the am battery case of the cylindrical battery (product number 18509), to obtain a negative electrode hop.

#### (d) Formation of Porous Film

[0335] The raw material paste for porous film was applied on both sides of the negative electrode hoop to give a thickness of 5 µm per side adhered to the negative electrode surface to form a porous film with varied surface roughness.

#### (e) Preparation of Electrolyte

[9336] For the non-aqueous electrolyte, a solvent mixture in which ethylene carbonate, ethyl methyl carbonate, and dimethyl carbonate were mixed with a volume ratio of 1:1:1, dissolving lithium hexafihorophosphate (LiPE<sub>d</sub>) to give a concentration of 1 mol/liter was used. Also, 3% by volume of vinylene carbonate relative to the solvent mixture was added to the non-succous electrolyte.

#### (f) Fabrication of Battery

[0337] The above positive electrode and negative electrode were cut to give a predestrained length, respectively, and were wound interposing a sheet separator with a thickness of 20 µm comprising a microprotus film made of polysthylene, and then were inserted into the battery case. Then, 5.5 g of the above electrolye was weighed, and into the battery case, and the opening of the case that the battery case with the polysthylene of the case of the control of the text of the control of the con

#### Battery G2

[0338] A battery G2 was made in the same manner as the battery G1, except that the viscosity of the raw material paste for porous lilm was controlled to be 92 cp.

#### Battery G3

[0339] A battery G3 was made in the same manner as the battery G1, except that the viscosity of the raw material paste for porous lilm was controlled to be 150 cp.

#### Battery G4

[0340] A battery G4 was made in the same manner as the battery G1, except that the viscosity of the raw material paste for porous film was controlled to be 182 cp.

#### Battery G:

[0341] A battery G5 was made in the same manner as the battery G1, except that the viscosity of the raw material paste for porous film was controlled to he 253 cp. Battery H1

[0342] A battery H1 was made in the same manner as the battery G1, except that the porous film was not formed on both sides of the negative electrode hoop.

## Battery H2

[0343] A battery I12 was made in the same manner as the battery G1. except that the viscosity of the raw material paste for porous film was controlled to be 17 cp.

#### Battery H3

[0344] A battery H3 was made in the same manner as the battery G1, except that the viscosity of the raw material paste for porous film was controlled to be 443 cp.

#### Battery G6

[0345] A battery G6 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the prorus film with a thickness of 0.3  $\mu m$  on both sides of the negative electrode.

#### Battery G7

[0346] A battery G7 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the porous film with a thickness of 0.5  $\mu m$  on both sides of the negative electrode.

#### Battery G

[0347] A battery G8 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the porous film with a thickness of 1 um on hoth sides of the negative electrode.

#### Battery G9

[0348] . A battery G9 was made in the same manner as the battery G3, except that the same raw material poste used in the battery G3 was used to form the porous film with a thickness of  $10~\mu m$  on both sides of the negative electrode.

## Battery G10

[0349] A battery G10 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the porous film with a thickness of 15 µm on both sides of the negative electrode.

# [0350] A battery G11 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the porous film with a thickness of 20 um on both sides of the negative electrode.

## Battery G12

[0351] A battery G12 was made in the same manner as the battery G3, except that the same raw material paste used in the battery G3 was used to form the porous film with thickness of 30 µm on both sides of the negative electrode.

## Battery G13

[0352] A battery G13 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 6 um.

## Battery G14

[0353] A battery G14 was made in the same manner as the hattery G3, except that the thickness of the sheet separator was set as 8 µm.

#### Battery G15

[0354] A battery G15 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 10 um.

#### Battery G16

[0355] A battery G16 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 15  $\mu$ m.

#### ..... C17

[0356] A battery G17 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 25 um.

## Battery G18

[0357] A battery G18 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 30 mm.

## Battery G19

[0358] A battery G19 was made in the same manner as the battery G3, except that the thickness of the sheet separator was set as 40 µm.

## Battery G20

[0359] A battery G20 was made in the same manner as the battery G3, except that the filler content relative to the total of the filler and the first binder was set as 30 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

#### Battery G21

[0360] A battery G21 was made in the same manner as the battery G3, except that the filter content relative to the total of the filter and the first binder was set as 50 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

#### Battery G22

[9361] A battery G22 was made in the same manner as the buttery G3, except that the filler content relative to the total of the filler and the first binder was set as 70 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

#### Battery G23

[9362] A battery G23 was made in the same manner as the battery G3, except that the filler content relative to the total of the filler and the first binder was set as 90 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

### Battery G24

[9363] A battery G24 was made in the same manner as the battery G3, except that the filler content relative to the total of the filler and the first binder was set as 95 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

## Battery G25

[0364] A battery G25 was made in the same manner as the battery G3, except that the filler content relative to the total of the filler and the first binder was set as 99 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

#### Battery 114

[0365] A battery H4 was made in the same manner as the battery G3, except that the filler content relative to the total of the filler and the first binder was set as 99.5 wt %, and a raw material paste with a viscosity of 150 ps at 20 rpm was prepared and used.

## Battery G26

[0366] A battery G26 was made in the same manner as the battery G3, except that water-soluble CMC was used instead of BM-72014 as a binder, and a raw material paste for porous film with a viscosity of 150 ps at 20 rpm was prepared and used. Herein, as a dispersion medium for the raw material paste, water was used instead of NMP.

#### Battery G27

[9367] A battery G27 was made in the same manner as the battery G3, except that water-insoluble PVDF (crystalline melting point of 174° C) was used instead of 18M-720H as a binder, and the raw material paste for porous film with a viscosity of 150 ps at 20 ym was prepared and used.

#### Battery H5

[0368] A battery 115 was made in the same manner as the battery G3, except that polyethylene beads with a median diameter of 0.3 µm was used instead of the alumina with a median diameter of 0.3 µm as the filler, and a raw material paste for porous film with a viscosity of 150 ps at 20 rpm was prepared and used.

#### Battery G28

[0369] A battery G28 was made in the same manner as the battery G3, except that titania with a median diameter of 0.3 pm as used instead of the alumina with a median diameter of 0.3 pm as the filler, and a raw material paste for porous film with a viscosity of 150 ps at 20 rpm was prepared and used

## Battery G29

[0370] In the fabrication of the negative electrode, instead of BM400B and CMC as a second binder, 8 wt % of PVDF. relative to the negative electrode active material was used. Herein, for PVDF, PVDF#1320 used for the binder of the positive electrode was used. Other than the above, a buttery G29 was made in the same manner as the battery G3.

#### . . .

[0371] The configuration of the porous film is summarized and shown in Table 11. Additionally, the thickness of the sheet separator and the kinds of the second binder used for the negative electrode are summarized and shown in Table

TABLE 11

				Pontus I	ilm		
						First Bin	Ser
	Paste	Film	Fill	ler		Crystalline Melting	Decomposition
Battery No	Viscosity (ps)	Thickness (jun)	Kind	Content (wt %)	Kind	Point (° C.)	Temperature (- C.)
Gt	48	5	Alumina	97	BM720H	None	320
G2	92	5	Alumina	97	BM720H	None	320

TABLE 11-continued

				Porous F	ilm		
						First Bind	ler
	Paste	Film	Fill	er		Crystalline Melting	Decomposition
Battery No.	Viscosity (ps)	Thickness (µm)	Kind	Content (wt %)	Kind	Point (* C.)	Temperature (° C.)
G3	150	5	Alumina	97	BM720H	None	320
G4	182	5	Alumma	97	BM720H	None	320
G5	253	5	Alumina	97	BM720H	None	320
G6	150	0.3	Alumma	97	BM720H	None	320
G7	150	0.5	Alumma	97	BM720H	None	320
G8	150	1	Alumens	97	BM720H	None	320
G9	150	10	Alumina	97	BM720H	None	320
G10	150	15	Alumina	97	BM720H	None	320
G11	150	20	Alumina	97	BM720H	None	320
G12	150	30	Alumina	97	BM720H	None	320
G13	150	5	Alamina	97	BM720H	None	320
G14	150	5	Alumina	97	BM720H	None	320
G15	150	5	Alumus	97	BM720H	None	320
G16	150	5	Alumma	97	BM720H	None	320
G17	150	5	Alumina	97	BM720H	None	320
G18	150	5	Alumina	97	BM720H	None	320
G19	150	5	Alumina	97	BM720H	None	320
G20	150	5	Alumma	30	BM720H	None	320
G21	150	5	Alumina	50	BM720H	None	320
G22	150	5	Alumina	70	BM720H	None	320
G23	150	5	Alumina	90	BM720H	None	320
G24	150	5	Alumina	95	BM720H	None	320
G25	150	5	Alumun	99	BM720H	None	320
G26	150	5	Ainma	97	CMC	None	245
G27	150	5	Alumina	97	PVDF	174	360
G28	150	5	Titanis	97	BM720H	None	320
G29	1.50	4	Alumina	97	BM720H	None	320
HI							
H2	17	5	Alumma	97	BM720H	None	320
113	443	5	Alumina	97	BM72011	None	320
H4	150	5	Alumina	99.5	BM720H	None	320
H5	150	5	PE beads	97	BM720H	None	320

[0372]

TABLE 12

			LI LINE				
			Porous	Film	Negative	Surface Roughness	
Battery No.	Separator	Negative		Flexibility	Flectrode		
	Thickness (jun)	Electrode Bunder	Adhesiveness	(No. of Defects)	Appearance No Change	Negative Electrode	Porous Film
G1	20	BM400B + CMC	OK	1	No Change	2.43	1.77
G2	20	BM400B + CMC	OK	0	No Change	2.43	0.83
G3	20	BM400B + CMC	OK	0	No Change	2.43	0.55
G4	20	BM400B + CMC	OK	0	No Change	2.43	0.96
G5	20	BM400B + CMC	OK	1	No Change	2.43	1.84
G6	20	BM400B + CMC	OK	0	No Change	2 43	2.21
G7	20	BM400B + CMC	OK	0	No Change	2.43	2.05
G8	20	BM400B + CMC	OK	0	No Change	2.43	1.31
G9	20	BM400B + CMC	OK	0	No Change	2.43	0.46
G10	20	BM400B + CMC	OK	0	No Change	2.43	0.40
G11	20	BM400B + CMC	OK	0	No Change	2.43	0.35
G12	20	BM400B + CMC	OK	1	No Change	2.43	0.41
G13	6	BM400B + CMC	OK	i)	No Change	2.43	0.55
G14	8	BM400B + CMC	OK	0	No Change	2.43	0.55
G15	10	BM400B + CMC	OK	0	Ne Change	2.43	0.55
G16	15	BM400B + CMC	OK	0	No Change	2.43	0.55
G17	25	BM400B + CMC	OK.	0	No Change	2.43	0.55

TABLE 12-continued

		Negative	Porous Film		Negative	Surface Roughness	
Bettery No.	Separator			Flexibility	Electrode	(um)	
	Thickness (µm)	Electrode Bander	Adhesiveness	(No. of Defects)	Appearance No Change	Negative Electrode	Porous Film
G18	30	BM400B + CMC	ок	0	No Change	2.43	0.55
G19	40	BM400B + CMC	OK	ō	No Change	2.43	0.55
G20	20	BM400B + CMC	OK	Ď.	No Change	2.43	0.81
G21	20	BM400B + CMC	OK	0	No Change	2.43	0.62
G22	20	BM400B + CMC	OK	0	No Change	2.43	0.60
G23	20	BM400B + CMC	OK	0	No Change	2.43	0.48
G24	20	BM400B + CMC	OK	0	No Change	2.43	0.43
628	20	BM400B + CMC	OK	ï	Changed	2.43	0.43
G26	20	BM400B + CMC	OK	4	No Change	2 43	0.57
G27	20	BM400B + CMC	OK	3	No Change	2 43	0.36
G28	20	BM400B + CMC	OK	ő	No Change	2.43	0.50
G29	20	PVDF	OK	2	No Change	2.28	0.52
H1	20	BM400B + CMC	774	6	No Change	2.43	0.32
H2	20	BM400B + CMC	OK	7	No Change	2.43	2.57
113	20	BM400B + CMC	OK	6	. to Change	2.43	3.61
H4	20	BM400B + CMC	NG	-	No Change	2.43	0.48
H.S	20	BM400B + CMC	ok	0	No Change	2.43	0.48

#### (Evaluation)

[Surface Roughness]

[0373] The surface roughness of the negative electrode surface before applying the porous film and the surface roughness of the porous film formed thereon after drying are measured respectively by a non-contact type surface roughness measuring instrument, to obtain an average roughness Ra. The results are shown in Table 12. [0374] Also, for the above produced porous film and the completed lithium ion secondary battery, porous film abssiveness, negative electrode appearance, porous film flexibility, battery designed capacity, charge and discharge characteristics of the battery, and the nail potentation assive a carderistics of the battery, and the nail potentation assive acventuated in the same manner as Example 2. The results are shown in Tables 12 to 13.

TABLE 13

						ail Penetr	stion Safety	
Battery No.	Designed Capacity (mAh)	Charge and Discharge Characteristics			Nail Speed 5 mm/s Temperature Reached		Nail Speed 180 mm/s Temperature Reached	
		Charge (mAiı)	400 mAh Discharge (mAh)	4000 mAh Discharge (mAh)	After 1 sec.	After 90 sec. (° C )	After 1 sec.	After 90 sec. (° C,)
Gl	1940	1933	1931	1829	75	89	76	91
G2	1936	1932	1930	1832	77	93	68	88
G3	1941	1940	1936	1847	74	94	fel	97
G4	1945	1941	1940	1842	72	92	77	94
G5	1939	1933	1931	1839	71	86	68	89
G6	2010	2011	2008	1887	68	123	68	89
G7	2011	2010	2006	1899	73	88	71	94
G8	2012	1999	1996	1901	70	89	74	93
G9	1875	1875	1874	1779	68	94	72	90
G10	1806	1799	1796	1708	71	95	73	87
G11	1735	1730	1728	1642	69	94	77	91
G12	1584	1581	1574	1386	71	96	72	88
G13	2157	2151	2146	2010	72	131	76	109
G14	2131	2124	2119	1989	73	87	71	90
G15	2093	2096	2086	1962	71	87	69	90
G16	2016	2016	2012	1902	68	95	67	88
G17	1874	1874	1870	1760	69	94	70	97
G18	1799	1797	1792	1679	72	88	68	88
G19	1656	1653	1649	1488	74	89	72	84
G20	1939	1938	1936	1491	73	91	71	85
G21	2014	1988	1961	1737	68	95	75	96

TABLE 13-continued

		Battery							
Battery No	Designed Capacity (mAh)				Nail Penetration Safety				
		Charge and Discharge Characteristics			Nail Speed 5 mm's Temperature Reached		Nail Speed 180 mm/s Temperature Reached		
		Charge (mAh)	400 mAh Discharge (mAh)	4000 mAh Discharge (mAh)	After 1 sec.	After 90 sec. (° C.)	After 1 sec.	After 90 sec. (° C.)	
G22	2015	1999	1989	1811	68	97	77	94	
G23	2015	2019	2015	1893	73	95	66	97	
G24	2015	2015	2009	1883	76	93	79	97	
G25	2017	2015	2010	1886	68	85	68	90	
G26	1941	1939	1938	1809	87	122	85	99	
G27	1943	1940	1939	1820	75	130	74	97	
G28	2016	2012	2005	1880	71	95	73	89	
G29	2014	1961	1958	1788	72	91	77	96	
H1	2014	2014	2003	1888	145		145		
H2	1944	1939	1936	1818	77	147	74	93	
H3	1938	1937	1931	1819	73	143	71	93	
H4	-	-						_	
H5	1944	1943	1939	1822	149	_	139	_	

#### (Results Consideration)

[0376] With regard to the thickness of the porous film, in the battery G12 having an excessively thick porous film, due to the shortened length of the electrode plate forming the electrode plate group, designed capacity decreased and the capacity at the high-rate discharge declined.

[0377] In the battery G19 in which the thickness of the separator was excessively large, due to the shortened length of the electrode plate forming the electrode plate group, the designed capacity decreased greatly and the capacity at the high-rate discharge declined.

[0.78] Then, in the battery (20 in which the filter content retaive to the total of the filter and the first bander was small (binder amount was large), the decline in the capacity at the high-rate discharge could be seen. This is probably beautions the gaps between the filter particles could not be obtained sufficiently due to the excessive first binder, to cannot be decline in ion conductivity of the porous film. As oppose to this, in the bettery H is which the amount of the binder sto cosmall, separations and chips of the porous film occurred frequently, due to the insufficient binding effects.

[0379]. In the battery G26 in which C/MC was used and the battery G27 in which PVDF was used as a first binder in the porous iflm, the heat generation could not be suppressed when the mail penetration speed was slowed. When these batteries were disassembled and checked, it was confirmed that the protus film also deformed, in addition to the separatior. [0380] Although in the battery G3 in which the binder had mibber elasticity, the porous film after winding kept its shape sufficiently and showed no defents, in the batteries G26 and G27, evaluation of flexibility for the porous film showed poor results. Further, in the battery G26, defective appearances due to the deformation of the negative electrode were seen after the formation of the proous film.

[0381] Also, in the battery G28 in which titania was used instead of alumina as a filler, it was confirmed that titania eachieved various functions same as alumina. On the other band, in the battery H5 in which polyethylene beads [P6] beads) were used, with regard to the nail penetration safety, it was the same level as the battery H1 having no porous

[0.882] Then, in the hattery (120 in which PVDF was used as the second binder, the binder content in the negative electrode became inevitably large, to cause a decrease in thitmin on acceptability of the negative electrode, and a gradual electrose in the churging capacity. Additionally, based on the properties of PVDF, the negative electrode plate hardward, thereby failing to make full use of the flexibility of the provise film.

[0383] Surface roughness of the porous film is discussed

[0384] Since all the negative electrodes used in this Esumple included a great amount of VGCF aiming an improvement of conductivity, surface roughness of the negative electrode was large. Separation of material mixture occurs easily in such negative electrode, due to lictions at which the raw material paste for protos film is centrolled in which the raw material paste for protos film is centrolled in which the raw material paste for protos film is centrolled in which the raw material paste for protos film is which the raw material paste for protos film is sledged on the negative electrode. Thus, this possible yields due to the separation of the material mixture is suppressed. In the hatterns of 10 of 28. surface roughness of the protoss film was regulated to become smaller than the surface roughness of the electrode plate. [0385] On the other hand, in the battery H2 in which the viscosity of the new material paste for protous film to be applied on the negative electrode was too small, the separation of the material mixture could not be avoided. This is because the asperities of the negative electrode surface were not smoothed by the perous film due to the run material paste with a large fluidity. Additionally, in the battery H3 in which the viscosity of the run material paste was excessively large, the surface roughness of the protous film necessary. The product of the process of the protous that the process of the protous film to the process of the protous film of the process of the pr

[9386] In the hattery II3 in which the separation of the material mixture did not occur, the acceleration for the material mixture did not occur, the acceleration for the generation could be seen in the nail penetration test as well. This is because uneven application occurred together with the increase in the surface roughness of the porous film. Since the unever application involves a partial void to porous film, the functions of the porous film to improve the safety is probably decreased.

[0387] Upon comparing the G6 and G7, it can be said that the average roughness Ra of helow 2.1 µm is desirable for the surface roughness of the porous film.

[0388] Herein, although the surface roughness of the purous film was changed based on the viscosity of the raw material paste, the optimal viscosity range easily changes depending upon the change in the composition of the porous film. Therefore, the viscosity range is not to be limited by this embodiness.

#### INDUSTRIAL APPLICABILITY

[0389] A lithium ion secondary battery of the present invention comprises a negative electroche having a high lithium ion acceptability, has a higher safety level, and can achieve a higher yield. Thus, the present invention is useful in the field which requires a higher safety and high-rate characteristic at the same time, especially as a power source for portable lectric devices and the like.

- A lithium ion secondary hattery comprising:
- a positive electrode capable of absorbing and desorbing lithium ion:
- a negative electrode capable of absorbing and desorbing lithium ion;
- a porous film interposed between said positive electrode and said negative electrode; and
- a non-aqueous electrolyte;
- wherein said porous film is adhered to a surface of at least said negative electrode,
- said porous film comprises an inorganic filler and a first binder, a content of said first binder in said porous film being 1.5 to 8 parts by weight per 100 parts by weight
- said first binder comprises a first rubber including an acrylonitrile unit, said first rubber heing water-insoluble and having a decomposition temperature of 250° C. or higher.

said negative electrode comprises a negative electrode active material capable of absorbing and desorbing lithium ion and a second binder, and

- said second binder includes a second rubber particle and a water-soluble polymer.
- The lithium ion secondary battery in accordance with claim 1, wherein said first rubber has a crystalline melting point of 250° C. or more.
- The lithium ion secondary battery in accordance with claim 1, wherein said first rubber includes a polyacrylomtrile chain.
- The lithium ion secondary battery in accordance with claim 1, wherein said first rubber comprises core-shell type
- particles and has an adhesive surface portion.

  5. The lithium ion secondary battery in accordance with claim 1, wherein said water-soluble polymer includes a
- methylcellulose unit.

  6. The lithium ion secondary battery in accordance with claim 1, wherein said second rubber particle includes a styrene unit and a butadiene unit.
- 7. The lithium ion secondary battery in accordance with claim 1, wherein a content of said second binder in said negative electrode is 1.5 to 3 parts by weight per 100 parts
- by weight of said negative electrode active material.

  8. The lithium ion secondary battery in accordance with claim I, wherein said inorganic filler comprises an inorganic oxide.

oxide.

9. The lithium ion secondary battery in accordance with claim 8, wherein a surface of said inorganic oxide is alkaline and has a BET specific surface area of 0.9 m<sup>2</sup>/g or more.

- 10. The lithium ion secondary battery in accordance with claim 1, wherein said inorganic oxide includes at least one selected from the group consisting of alumina and titanium oxide.
- 11. The lithium ion secondary battery in accordance with claim I, wherein a surface roughness of said porous film is less than a surface roughness of an electrode surface to which said porous film is adhered to.

12. The lithium ion secondary battery in accordance with claim I, wherein said inorganic filler comprises a mixture of a large particle group and a small particle group, and an average particle size A of said large particle group and an average particle size B of said small particle group satisfy the formula (1):

#### 0.055R(A.50.2

- 13. The lithium ion secondary battery in accordance with claim 1, wherein said positive electrode and said negative electrode are wound with said porous film interposed therefore.
- 14. The lithium ion secondary battery in accordance with claim 1, wherein a thickness of said porous film is 0.5 µm or more and 20 µm or less.
- 15. The lithium ion secondary battery in accordance with claim 1, wherein a separator is further interposed between said positive electrode and said negative electrode.
- 16. The lithium ion secondary battery in accordance with claim 15, wherein a thickness of said separator is 8 μm or more and 30 μm or less.

. . . . .

# **Polyacrylonitrile**

From Wikipedia, the free encyclopedia

Polyacrylonitrile (PAN) is a resinous, fibrous, or rubbery organic polymer. Almost all polyacrylonitrile resins are copolymers made from mixtures of monomers with aerylonitrile as the main component. PAN fibers are the chemical precursor of high-quality carbon fiber. It is chemically modified to make the carbon fibers found in plenty of both high-tech and common daily applications such as civil and military aircraft primary and secondary structures, missiles, solid propellant rocket motors, pressure vessels, fishing rods, tennis rackets, badmitton rackets & high-tech bicycles. It is a component repeat unit in several important copolymers, such as styrene-acrylonitrile (SAN) and aerylonitrile butadeine styrene (ABS) plastic.



Chemical structure of the repeating unit of polyacrylonitrile

## **Applications**

Homopolymers of polyacrylonitrile have been used as fibers in hot gas filtration systems, outdoor awnings, sails for yachts, and even fiber reinforced concrete. Copolymers containing polyacrylonitrile are often used as fibers to make knitted clothing, like socks and sweaters, as well as outdoor products like tents and similar items. If the label of a piece of clothing says "acrylic", then it is made out of some copolymer of polyacrylonitrile. It was made into spun fiber at DuPont in 1941 and marketed under the name of Orlon. Acrylonitrile is commonly employed as a comonomer with styrene (e.g. SAN, ABS, and ASA (acrylonitrile styrene acrylate) plastics).

### External links

Polyacrylonitrile at Polymer Science Learning Center

Retrieved from "http://en.wikipedia.org/wiki/Polyacrylonitrile"

Categories: Polyacrylates | Copolymers | Synthetic resins | Synthetic fibers | Thermoplastics | Polymer stubs

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APPENDIX 3

# Polyacrylic acid

From Wikipedia, the free encyclopedia

Poly(acrylic acid) or PAA or Carbomer is a type of anionic polymer. The monomer of poly(acrylic acid) is acrylic acid. In a water solution at neutral pH, many of the side chains of PAA will lose their protons and acquire a negative charge. This makes PAA a polyclectrolyte.

Dry PAA is a white solid. It is capable of adsorbing many times its weight in water, and hence is used in disposable diapers[1].

It also is used as a thickening agent. It inactivates Chlorhexidine gluconate.

## See also

- Sodium polyacrylate
- Carbomer

### References

http://pslc.ws/macrog/acrylate.htm

Retrieved from

"http://en.wikipedia.org/wiki/Polyacrylic acid" Categories: Polymers | Polyacrylates | Polymer stubs Poly(acrylic acid)

Identifiers

9003-01

Properties Molecular formula  $(C_3H_4O_2)_n$ Molar mass

### Hazards

EU classification

variable

R-phrases

R36 R37 R38

√(what is this?) (verify) Except where noted otherwise, data are given for materials in their standard state (at 25 °C, 100 kPa)

Infobox references

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# Acrylate polymer

From Wikipedia, the free encyclopedia (Redirected from Polyacrylate)

An acrylate polymer belongs to a group of polymers which could be referred to generally as plastics. They are noted for their transparency and resistance to breakage and elasticity. Also commonly known as acrylics or polyacrylates.

Typical acrylate monomers used to form acrylate polymers are: acrylic acid, methyl methacrylate and acrylonitrile.

### Contents

- 1 Monomers
- 2 Acrylic elastomers
- 3 Other acrylic polymers
- 4 References
- 5 See also

## Monomers

Other examples of acrylate monomers are:

- Methacrylates
- · methyl acrylate
- · ethyl acrylate
- 2-chloroethyl vinyl ether
- 2-ethylhexyl acrylate.
- hydroxyethyl methacrylate
- butyl acrylate
- butyl methacrylate
- TMPTA.

## Acrylic elastomers

Acrylic elastomer is a general term for a type of synthetic rubber whose main component is acrylic acid alkylester (ethyl or butyl ester). [1] Acrylic elastomer has characteristics of heat and oil resistance.

It is divided into old type and new type: Old types include ACM (copolymer of acrylic acid ester and 2chloroethyl vinyl ether) containing chlorine and ANM (copolymer of acrylic acid ester and acrylonitrile) without chloride. Other than the slightly better water resistance of ANM, there are no physical differences; even processability is poor for both types. Since prices are also high, demand is not so high vis-à-vis the characteristics. On the other hand, the new type of acrylic rubber does not contain any chlorine despite its unclear chemical composition. Processability has been improved, and most of tackiness to rolls as well as staining problems related to molds have been solved.

Major characteristics of acrylic rubber include heat resistance and oil resistance; it can endure a temperature of  $170 \sim 180^{\circ}$ C under dry heat or in oil. Since it does not have a double bond, acrylic rubber

also boasts of good weatherability and ozone resistance.

The materials are used mainly for oil seals and packagings related to automobiles.

## Other acrylic polymers

- Polymethyl methacrylate, an acrylate polymer familiar to consumers is the clear break resistant glass or sheeting sold in hardware stores as acrylic glass or under the trade name Plexiclas.
- Polyacrylate emulsion, water-born coating, are used as binder for outdoor and indoor "latex" house paints
- Acrylic paints as artist paints
- Acrylic fibre
- Sodium polyacrylate water soluble thickeners, a polymer for the production of the Superabsorbent polymer (SAP) used in disposable diapers due to its high absorbency per unit mass
- Acrylic resin as pressure-sensitive adhesive
- "Super glue" is a formulation of cyanoacrylate.
- PVAc copolymer emulsion adhesive of vinyl acetate (VAM) and acrylic acid (VAA)
- Polyacrylamide copolymer used as floculation agent in water treatment

## References

1. http://www.patentstorm.us/patents/6015860/claims.html

## See also

- (Meth)acrylates
- · Acrylic (disambiguation)

Retrieved from "http://en.wikipedia.org/wiki/Acrylate\_polymer"

Categories: Polyacrylates | Plastics | Optical materials

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organization.



Keywords gel

Acrylates are a family of polymers, which are a type of vinyl polymer.
Acrylates are of course made from acrylate monomers, and it's about time we explained what those are. Acrylate monomers are esters which contain vinyl groups, that is, two carbon atoms double-bonded to each other, directly attached to the carbonyl carbon.

Some acrylates have an extra methyl group attached to the alpha carbon, and these are called methacrylates. One of the most common methacrylate polymers is poly(methyl methacrylate).

H C=O In case you've never heard of the element R, there is none. R is a wild card, and it stands for any atom or group af atoms.

polymerization of methyl methacrylate to get poly(methyl methacrylate)

### Acrylate and Methacrylate

#### one of Nietzsche's lesser known works

One might not think that this little methyl group would make a whole lot of difference in the behavior and properties of the polymer, but it does. Poly(methyl acrylate) is a white rubber at room temperature, but poly(methyl methacrylate) is a strong, hard, and clear plastic.

This is poly(methyl acrylate). This is poly(methyl methacrylate). It is soft and rubbery. It is a hard plastic.

As it turns out, how soft or hard a polymer is at a given temperature is determined by what we call chain mobility, that is, how well the polymer chains wiggle past and around each other. The more they can move, the softer the polymer is. It helps here to think the scene in the movie Raiders of the Lost Ark, in which our hero Indiana Jones finds himself in an Egyptian temple and ankle deep in slithery poisonous snakes. Polymer chains are like those snakes. Smooth snakes can move past each other very easily. But if you could imagine those snakes having giant barbed spikes all up and down their backs like some of the dinosaurs in another Steven Spielberg film, you can see that they might not be moving around quite as much on the floor of that Egyptian temple. Their spikes would catch on each other, and slithering would become quite difficult.

Poly(methyl methacrylate) is like one of those snakes with giant barbed spikes all up and down its back, with those extra methyl groups acting like the spikes to put a quick stop to any slithering the poly(methyl methacrylate) chains would try to do. Poly(methyl acrylate), on the other hand, is like the smooth snakes. Without that extra methyl group getting in the way, they can slither all they want. If the polymer chains can slither and wiggle past and around each other easily, the whole mass of them will be able to flow more easily. So, a polymer which can move around easily will be soft, and one which can't will be hard, to put it simply.

If you want to know more about how polymers are like snakes take a look at the glass transition temperature page.

### No More Saggy Diapers That Leak

The simplest acrylate polymer is one of the least well understood. And that would be...

### Poly(acrylic acid)!

This is what we call a polyelectrolyte. That is to say, each repeat unit has an ionizable group. In this case, it's a carboxylic acid group. Poly(acrylic acid) is bizarre because it soaks up water like crazy. It absorbs many times its own weight in water with no problem. Polymers that do this are called superabsorbers. So some bright individual had the bright idea of putting this stuff into baby diapers. In a diaper poly(acrylic acid) absorbs the little mess your little bundle of joy leaves behind. Remember all those diaper commercials where they pour that blue stuff on the diapers and it disappears? What you saw was poly(acrylic acid) in the act of superabsorption.

And don't think for one minute that we have the slightest idea why poly(acrylic acid) can absorb so much water. We're completely cluckes!

The advantage of having diapers with poly(acrylic acid) in them, aside from the fact that they're less messy, is that once the mess is locked up in the poly(acrylic acid), baby doesn't have to sit in it until mom and dad figure out that it's time for a change. Otherwise, baby could get some unpleasant skin rashes.

## A Little Nitrogen Music

There are several derivatives of polyacrylates which contain nitrogen. Polyacrylamide and polyacrylonitrile are two shown in the picture. Polyacrylonitrile is used to make acrylic fibers. Polyacrylamide will dissolve in water and is used industrially. Crosslinked polyacrylamides can absorb water. (Crosslinked polymers can't really dissolve, if you think about it.) These gels are used to make soft contact lenses. It's the absorbed water in them that makes them soft. To see the polymer and its monomer acrylamide in 3-D, click here.

$$\begin{array}{cccc} H & H & \\ C = C & \longrightarrow & -tCH_2 - CH_{\frac{1}{2}} \\ K & C = C & & C = C \\ K & KH_2 & & KH_2 \\ \end{array}$$

acrylamide and polyacrylamide

$$\begin{array}{c} H \\ C = C \\ H \end{array} \longrightarrow \begin{array}{c} - t \operatorname{CH}_2 - \operatorname{CH}_n \\ C \equiv N \end{array}$$

acrylonitrile and polyacrylonitrile

Now read a story I wrote about acrylates. It's called...

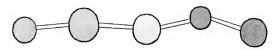
What Wonders Acrylates Can Do.



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translations

Wikipedia

Derivative (chemistry)

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example

In chemistry, a derivative is a compound that is derived from a similar compound or a compound that can be imagined to arise from another compound, if one atom is replaced with another atom or group of atoms. [1] The latter definition is common in organic chemistry. In biochemistry, the word is used about compounds that at least theoretically can be formed from the precursor compound. [2]

wikipedia

Chemical derivatives may be used to facilitate analysis. For example, melting point (MP) analysis can assist in identification of many organic compounds. A crystalline derivative may be prepared, such as a semicarbazone or 2,4-dinitrophenylhydrazone (derived from aldehydes/ketones), as a simple way of verifying the identity of the original compound, assuming that a table of derivative MP values is available. (3) Prior to the advent of spectroscopic analysis, such methods were widely used.

### See also

Derivatization

#### References

- 1. ^ "Definition of Derivative". Chemicool. 2007-09-18.
- http://www.chemicool.com/definition/derivative.html. Retrieved 2007-09-18.
- Oxford Dictionary of Biochemistry and Molecular Biology. Oxford University Press. ISBN 0-19-850673-2.
- \* Williamson, Kenneth L. (1999). Macroscale and Microscale Organic Experiments, 3rd ed.. Boston: Houghton-Mifflin. pp. 426-7. ISBN 0-395-90220-7.

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→ All translations of DERIVATIVE CHEMISTRY

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APPENDIX 7



- · Periodic Table
- · Elements A to Z

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## **Definition of Derivative**

A compound that can be imagined to arise from a partent compound by replacement of one atom with another atom or group of atoms. Used extensively in orgainic chemistry to assist in identifying compounds.

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APPENDIX 8

HOMEBROWSEWORKSPACESHOPBLOGADVOCACYHELPLOG IN Reset Submit Query Home Search Site Encyclopædia Britannica Table of Contents ▼ Article Related Articles Citations Related Articles. Ebooks & More EDIT SAVE PRINT F-MAII Web Links Article History carboxylic acid derivative Contributors LINKS Related Articles Get involved Share Dictionary & Thesaurus Aspects of the topic carboxylic acid Widgets derivative are discussed in the following places at Britannica Assorted References · major reference (in carboxylic acid (chemical compound): Derivatives of carboxylic acids) The carboxylic acid derivatives discussed here (with the exception of nitriles) share the RCO structure with aldehydes, ketones, and carboxylic acids themselves · carboxylic acids (in carboxylic acid (chemical compound)) Compounds in which the -OH of the carboxyl group is replaced by certain other groups are

Compounds in which the "OH of the carboxyl group is replaced by certain other groups are called carboxylic acid derivatives, the most important of which are acyl halides, acid anhydrides, esters, and amides

functional groups (in chemical compound: Carboxylic acids)

The structural unit containing an alkyl group bonded to a carbonyl group is known as an acyl group. A family of functional groups known as carboxylic acid derivatives contains the acyl group bonded to different substituents

#### Other

The following is a selection of items (artistic styles or groups, constructions, events, fictional characters, organizations, publications) associated with "carboxylic acid derivative"

- · acid halide (chemical compound)
- · carbonate (chemical compound)
- · carboxylic acid (chemical compound)